

The genus Andrena Fabricius, 1775 in the Iberian Peninsula (Hymenoptera, Andrenidae)

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Abstract

The Iberian Peninsula is a global hotspot for bee diversity due to its large number of different habitats, particularly Mediterranean scrubland, mountains, and hot and cold steppe. In line with its status as a hotspot of bee diversity, the peninsula hosts a very large Andrena fauna, which despite progress in recent years remains incompletely studied, particularly with reference to genetic investigation. Here the Iberian Andrena fauna is comprehensively revised, resulting in a total of 228 recorded species. Numerous taxonomic changes are necessary following inspection of museum specimens, type material, and genetic investigation. The following subgenera are described: Pruinosandrena subgen. nov., containing six taxa previously placed in the subgenus Campylogaster Dours, 1873, and Blandandrena subgen. nov., Bryandrena subgen. nov., Limbandrena subgen. nov., and Ovandrena subgen. nov., containing one, one, one, and four taxa previously placed in the subgenus Poliandrena Warncke, 1968. Andrena (Limbandrena) toelgiana Friese, 1921 **syn. nov.** is synonymised with A. (Limbandrena) limbata Eversmann, 1852. The current lectotype of A. (Micrandrena) obsoleta Pérez, 1895 was incorrectly designated by Warncke; the taxon differs from A. obsoleta sensu Warncke, belonging instead to a taxon within the A. mariana Warncke, 1968 complex. A new lectotype is designated for A. obsoleta sp. resurr. from Algeria, and A. mariana solda Warncke, 1974 syn. nov. is synonymised with it; A. (Micrandrena) alma Warncke, 1975 stat. nov., A. (Micrandrena) mica Warncke, 1974 stat. nov., and A. (Micrandrena) tenostra Warncke, 1975 stat. nov. are raised to species status. Andrena (Truncandrena) abunda Warncke, 1974 stat. nov., A. (Micrandrena) lecana Warncke, 1975 stat. nov., A. (Pruinosandrena) parata Warncke, 1967 stat. nov., A. (Micrandrena) pauxilla Stöckhert, 1935 sp. resurr., A. (Pruinosandrena) succinea Dours, 1872 sp. resurr., and A. (Notandrena) varuga Warncke, 1975 **stat. nov.** are also returned or elevated to species status. A lectotype is designated for A. (Euandrena) lavandulae Pérez, 1902 sp. resurr. which is returned to species status, and A. (Euandrena) impressa

Warncke, 1967 syn. nov. is synonymised with it. Andrena (Truncandrena) nigropilosa Warncke, 1967 stat. **nov.** is elevated to species status, and A. (Truncandrena) truncatilabris espanola Warncke, 1967 syn. nov. is synonymised with it as a junior subjective synonym. A lectotype is designated for A. (Melandrena) vachali Pérez, 1895; A. (Melandrena) creberrima Pérez, 1895 syn. nov. and A. (Melandrena) vachali syn. nov. are synonymised with A. (Melandrena) discors Erichson, 1841, and Andrena (Melandrena) hispania Warncke, 1967 **syn. nov.** is synonymised with A. (Melandrena) morio Brullé, 1832. Andrena (Pruinosandrena) mayeti Pérez, 1895 syn. nov. is newly synonymised with A. (Pruinosandrena) caroli Pérez, 1895 and A. (incertae sedis) setosa Pérez, 1903 syn. nov. is newly synonymised with A. (incertae sedis) ranunculorum Morawitz, 1877. Andrena (Simandrena) cilissaeformis Pérez, 1895 sp. resurr. is returned to species status, and is the correct name for A. (Simandrena) breviscopa auctorum. Andrena (incertae sedis) breviscopa Pérez, 1895 is returned to synonymy with A. (incertae sedis) numida Lepeletier, 1841, and A. (incertae sedis) inconspicua Morawitz, 1871 is newly synonymised syn. nov. with A. numida. Andrena (Euandrena) isolata sp. nov. and A. (Micrandrena) ortizi sp. nov. are described from the Sierra Nevada (Granada), A. (Truncandrena) ghisbaini sp. nov. is described from Málaga province, and A. (Avandrena) juliae sp. nov. is described from Cádiz province. The males of A. (Micrandrena) alma and A. (?Euandrena) ramosa Wood, 2022 are described. Additional lectotypes are designated for A. (Plastandrena) asperrima Pérez, 1895, A. (Plastandrena) atricapilla Pérez, 1895, A. (Aenandrena) hystrix Schmiedeknecht, 1883, A. (Pruinosandrena) lanuginosa Spinola, 1843, A. (Notandrena) ranunculi Schmiedeknecht, 1883, and A. (Euandrena) symphyti Schmiedeknecht, 1883. Neotypes are designated for A. (Chlorandrena) boyerella Dours, 1872, A. (Notandrena) griseobalteata Dours, 1872, A. (Taeniandrena) poupillieri Dours, 1872, A. (Pruinosandrena) succinea Dours, 1872, and A. (incertae sedis) numida Lepeletier, 1841. Type photographs and diagnostic characters are presented in each case, as well as new dietary information for understudied species. Finally, an identification key is presented in order to facilitate future research on this hyper-diverse genus in one of their global diversity hotspots, and current and future research perspectives for Iberian Andrena are discussed.

Keywords

Cryptic species, DNA barcoding, Iberian endemic species, solitary bees, taxonomy

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Introduction

The Iberian Peninsula is one of the best places to find and study wild bees globally, with more than 1,000 species present due to its dry and warm climate, abundance of open seasonal habitats, status as a glacial refugium, and rich diversity of flowering plants (Lhomme et al. 2020; Ortiz-Sánchez 2020; Orr et al. 2021). Within this large fauna, the most speciose element consists of bees from the genus Andrena which are abundant throughout the peninsula and which are a ubiquitous component of the spring pollinator community. This pattern is typical for the Holarctic, as Andrena are the second largest genus of bees globally with around 1,650 species following recent revisions (Ascher and Pickering 2020; Pisanty et al. 2022a; Wood and Monfared 2022) and are almost always the most speciose genus in regional faunas across this region. Considering that bees emerged in the early to mid-Cretaceous period 110-140 million years ago (Danforth et al. 2013; Branstetter et al. 2017; Murray et al. 2018), as a major bee genus Andrena is relatively young, arising around 25 million years ago (Pisanty et al. 2022b). Given the size of the genus, Andrena has speciated extremely quickly (Bossert et al. 2022), leading to enormous species richness and often extreme taxonomic difficulty in delineating and recognising species. Given the abundance and ubiquity of this genus in the Iberian fauna, it is desirable to have suitable taxonomic resources to enable study of this rich fauna. However, nothing resembling an accessible revision is currently available.

Despite the great number of species present in Iberia, very few authors have worked on this fauna in any detail, certainly compared to that of north-western Africa (e.g. Erichson 1841; Lepeletier 1841; Dours 1872, 1873; Pérez 1895, 1902, 1903; Morice 1899; Schmiedeknecht 1900; Saunders 1908; Benoist 1961). Somewhat incredibly, before 2020, only six papers were published that described currently valid Andrena species with a locus typicus in Iberia: Erichson (1835, three species), Dours (1873, one species), Pérez (1895, six species), Pérez (1902, one species), Warncke (1967, 10 species), and Warncke (1975a, 15 species), although some other works described what are now synonymous names (e.g. Spinola 1843; Pérez 1903; Friese 1922). The fauna was therefore quite unstudied before the revisionary work of Warncke (1967, 1975a, 1976), and following these publications almost no additional taxonomic work was carried out on the Iberian fauna until very recently. This attention has come in the form of revisionary taxonomic work and faunal work that has provided new distributional records, demonstrated the presence of doubtful species in Iberia, and recorded new species for the Iberian Andrena fauna (Baldock et al. 2018; Álvarez Fidalgo et al. 2020, 2021a, 2021b, 2022a, 2022b; Wood et al. 2020a, 2021, 2022; Alvarez Fidalgo and Aguado Martín 2022; Wood 2022; Wood and Ortiz-Sánchez 2022).

As a result of these collective efforts, our understanding of Iberian *Andrena* is as great as it has ever been. However, commencing ecological or taxonomic work on this fauna remains highly challenging due to the lack of accessible identification resources. Valuable information is available in original descriptions, group revisions (e.g. Schwenninger 2015), and treatments on other regional faunas (e.g. Schmid-Egger and Scheuchl 1997; Amiet et al. 2010), as well as in the global revisionary catalogue of Gusenleitner and Schwarz (2002). However, digging through this mountain of infor-

mation is time consuming and sometimes dispiriting and confusing, in part due to persistent problems with species concepts and taxonomic nomenclature. It is therefore the objective of this present paper to resolve most of the outstanding taxonomic problems with the Iberian *Andrena* fauna and to present a new synthesis of this information in the form of a comprehensive identification key. Due to the strong faunal links between North Africa and Iberia, it is necessary to revise parts of the North African *Andrena* fauna in parallel; a subsequent paper dealing with taxonomic revisions exclusive to the North African *Andrena* fauna will follow shortly. Therefore, several taxonomic changes will be made here that affect the North African fauna, although this is not the principal objective of the current work. Finally, novel ecological data are provided concerning the pollen foraging niche of unstudied Iberian or West Mediterranean *Andrena* species.

Methodology

Species concepts

It is important to briefly discuss species concepts in the context of this work. Because *Andrena* taxonomy in the West Palaearctic region was dominated by Klaus Warncke in the second half of the 20th Century, it is his classification system that has largely been followed by subsequent workers (e.g. Gusenleitner and Schwarz 2002). Warncke recognised species solely based on their morphology, and used subspecies to delineate geographical variation. Subsequent workers have treated many of these subspecies as valid species based on morphological, genetic, and ecological data (e.g. Gusenleitner and Schwarz 2002; Schmid-Egger 2005; Schwenninger 2015; Praz et al. 2019; Kratochwil 2021; Wood et al. 2021; McLaughlin et al. 2022; Pisanty et al. 2022a).

To date, there has not been a deep discussion of species concepts in *Andrena* compared to better studied bee groups such as bumble bees (*Bombus*, e.g. Williams et al. 2020; Rasmont et al. 2021). Use of a purely biological species concept in *Andrena* is premature in almost all cases as we have extremely limited understanding of their mating systems and pre- and post-zygotic reproductive barriers, with this only being estimated *post hoc* by quantifying realised gene flow (e.g. McLaughlin et al. 2022). Given that the study of *Andrena* taxonomy using genetics is still in its infancy, it is best to consider *Andrena* species to be evolutionarily independent lineages (de Queiroz 2007), and that through the application of integrative taxonomy (including genetic, morphological, and ecological data), one can arrive at a robust species concept (Schlick-Steiner et al. 2010).

As such, the present work builds on the morphological species concepts developed by previous workers and integrates genetic and ecological lines of evidence in order to ensure evidence-based species delineation. In this context, subspecies are used pragmatically, following the position of Mayr (1963, see Rasmont et al. 2021) and drawing from the taxonomic heritage of Warncke's West Palaearctic revisions. Where sufficient data are available, these subspecies are validated as species in their own right or synonymised. Where such data are not available or the results are ambiguous, subspecies are retained in order to indicate future avenues of taxonomic study.

Genetic sampling and analysis

Andrena specimens were sampled in Iberia and Morocco, predominantly during May–July 2021 in Spain and March–July 2022 in Morocco, but also using specimens collected in previous years. For genetic barcoding, a single midleg was removed from pinned specimens and sent to the Canadian Center for DNA barcoding (CCDB) in Guelph, Canada, for DNA extraction and sequencing (Ivanova et al. 2006). Specimens were sequenced following standardised high-throughput protocols. Both Lep1 and BeeCox1F1 primers were used (Hebert et al. 2004; Bleidorn and Henze 2021) to target the COI-5 region. All sequences are published on the Barcode of Life Database (BOLD) website under the public dataset "DS-ANDWMED".

Phylogenetic trees were supplemented with additional published sequences (e.g. Schmidt et al. 2015) that were downloaded from Genbank and the Barcode of Life Data System. Trees were also significantly enriched with sequences produced by the Centro de Investigação em Biodiversidade e Recursos Genéticos (CIBIO, Portugal) as part of a collaboration to barcode the Portuguese bee fauna that will be published in the near future, and sequences will be available on BOLD. Sequences beginning with the code 'IBIHM' were generated by CIBIO. Sequences were aligned using MAFFT (Katoh and Standley 2013). Aligned sequences were analysed in Seaview (Gouy et al. 2010) using a maximum likelihood analysis which was run with 1000 bootstraps. Intra- and interspecific distances were calculated using MEGA-X (Kumar et al. 2018). Outgroups were chosen based on the subgeneric analysis of Pisanty et al. (2022b) in order to ensure that the selected outgroup species is part of a subgenus or species group more basal to those selected for focused study.

Taxonomic decisions were informed by analysis of the COI gene. However, decisions were not taken exclusively on the basis of these analyses, as there are a number of inherent limitations when using this fragment to inform taxonomy. The COI fragment itself only represents a single locus of mitochondrial DNA which is inherited matrilineally, can introgress across species boundaries, can transfer to the nuclear genome, cannot detect hybridisation, and can produce topologies that do not represent species trees (e.g. Toews and Brelsford 2012). Species delineation based solely on differences in mitochondrial DNA is therefore discouraged, with integrative taxonomic preferred (Padial et al. 2010; Gallego-García et al. 2023). More broadly, more powerful genetic techniques are now available such as Ultra Conserved Elements that allow the generation of sequences from thousands of loci. These have been applied to bee taxonomy at the species-level, and whilst they often produce results concordant with COI analysis (e.g. de Oliveira Andrade et al. 2022; McLaughlin et al. 2022), they can also clearly resolve problems for which COI analysis produces ambiguous or inconclusive results (e.g. Gueuning et al. 2020). However, to date, UCEs have predominantly been employed to resolve specific questions, and not to sample across a fauna in order to have the broadest possible taxonomic coverage, as was the objective in this study. Given these limitations, combined with the conceptual points discussed above, analysis of the COI fragment is used as an additional line of evidence to support taxonomic decision making, and is never used as the sole metric.

More broadly, COI analysis is most useful when dealing with species-specific alpha taxonomic decisions. Due to the rate at which this fragment accumulates mutations, whilst closely related species are grouped together, more distantly related species often show no clear structuring and the true evolutionary relationships are not captured. For this, more conserved genes must be sampled, such as by using UCEs. In this work, several new *Andrena* subgenera are described and illustrative phylogenetic trees are presented based on analysis of the COI fragment; the description of these subgenera is based on the work of Pisanty et al. (2022b) who identified undescribed clades of *Andrena*. These clades were not discovered here through analysis of the COI fragment, and these illustrative phylogenetic trees should not be considered strong support for their existence.

Finally, there are several cases presented here where morphological and genetic data produce ambiguous results concerning the status of certain taxa. Several of these taxa are widespread, and are usually described from outside of the Iberian Peninsula (e.g. see problems with paraphyly of *A. hedikae* Jäger, 1934). In these cases, no taxonomic action is taken, as it is preferable to have topotypic genetic samples and to consider these taxa across their entire nominal range. Where multiple valid species are potentially present, these species are referred to using the phrase 'aggregate' to reflect this situation. In contrast, when species are described from Iberia or have ranges that are restricted to the peninsula or to the West Mediterranean region, taxonomic decisions can be and are made with a greater degree of confidence which reflects the stronger and more complete evidence base available here.

Checklist and identification

For updating the Iberian *Andrena* species total, the checklist of Ortiz-Sánchez (2020) for mainland Spain and Portugal is used as a baseline. As this list is mostly correct, it is not considered necessary to produce a full annotated list for the Iberia *Andrena* fauna here; instead, a detailed justification will be given for the changes which have been made since 2020 or which are newly made here. A full checklist is provided in Suppl. material 1.

Identification key and geographic scope

For the identification keys, the female key is partly based on an unpublished key to Iberian *Andrena* written by Klaus Warncke (in German) that was kindly shared with me by Erwin Scheuchl (Ergolding, Germany). This key contained around 170 species, so considerable modifications were needed to account for the substantially larger faunal total recorded here, as well as accounting for newly described species and other taxonomic changes. The male key is novel, but both the female and male keys have been strongly inspired by the keys of Schmid-Egger and Scheuchl (1997) and Amiet et al. (2010). Additional characters have also been integrated from published works such as Schönitzer et al. (1995), Schwenninger (2009, 2013, 2015), and Praz

et al. (2019, 2022). It is not always easy to remember the precise source of specific characters used here that have been accumulated and integrated over several years of study, as many are scattered throughout the literature in original descriptions, some derive from my own observations, and many are taken from the important diagnostic characters that are embedded throughout Gusenleitner and Schwarz (2002), but which can be somewhat obscure and hidden in individual species accounts. It is my hope that this key can bring this information together as a novel synthesis, becoming a more easily digestible resource.

The geographic scope of the key is limited to the Iberian Peninsula. It cannot be used in North Africa due to the many different or endemic faunal elements found there; for example, it only covers 114 of the 201 (56.7%) Andrena species known from Morocco (Wood in prep.). In a European context, the key can be used in the Balearic Islands, though only a fraction of the species covered in this work occur there. It can be generally used in southern France up to (but not including) the Maritime Alps, but some taxa are missing, such as endemic species (e.g. Andrena (Taeniandrena) vocifera Warncke, 1975) or widespread European species that do not cross the Pyrenees into Iberia (e.g. Andrena (Micrandrena) pusilla Pérez, 1903). However, the existing keys of Schmid-Egger and Scheuchl (1997) and Amiet et al. (2010) can be used in conjunction with this Iberian key to cover the vast majority of the Andrena fauna of southern France. The Iberian key should not be used in an Italian context as the Italian fauna is already sufficiently distinct to render the key of limited use, either in northern Italy due to the presence of eastern faunal elements (e.g. Andrena (Aenandrena) bisulcata Morawitz, 1877) or in Sicily due to the large number of non-Iberian North African elements present there, and also because many Iberian taxa are absent from the Italian fauna which has a much lower degree of endemism. Finally, this key should not be used in Corsica or Sardinia, as their island faunas require dedicated study due to the presence of endemic and North African species, as well as local forms or subspecies.

Distribution maps

No distribution maps are presented as part of this work, as they are for other revisions such as that of Ortiz-Sánchez and Pauly (2017). This is because whilst the material that I have examined, validated, and digitised is sufficient to allow a more or less complete taxonomic understanding of the Iberian fauna, it is not sufficiently geographically comprehensive, and hence distribution maps would be incomplete at the scale of the peninsula. Moreover, the distribution maps of Warncke presented by Gusenleitner and Schwarz (2002) are sufficiently correct to be informative to students of the Iberian fauna, with the assumption that the updated taxonomy presented here and by other workers after 1993 is integrated. Important distributional information is given in the text where relevant, and also in the identification key, as many taxa (particularly Euro-Siberian taxa) are geographically limited and the place of capture can strongly inform their identification (e.g. species restricted in an Iberian context to the Pyrenees, to central or southern Spain, etc).

Dietary niches of Iberian Andrena species

Pollen was removed from female *Andrena* specimens in order to quantify the pollen foraging niche of understudied species. Specimens were selected from Iberia and other Mediterranean countries when the species' range extends beyond the peninsula. Pollen was removed, processed, and identified following the methodology of Wood and Roberts (2018). Dietary classification (polylecty, mesolecty, oligolecty, narrow oligolecty) follows Müller and Kuhlmann (2008).

Morphological terminology

Morphological terminology follows Michener (2007). Specimens were measured from the centre of the clypeus at the front of the head to the apical tip of the metasoma to the nearest 0.5 mm. Photographs were taken using an Olympus E-M1 Mark II with a 60 mm macro lens. Additional close-ups were taken with the addition of a Mitutoyo M Plan Apo 5X infinity corrected objective lens. Photographs were stacked using Helicon Focus B (HeliconSoft, Ukraine) and plates were prepared in GNU Image Manipulation Program (GIMP) 2.10. Post-processing of some images was made in Photoshop Elements (Adobe Systems, USA) in order to improve lighting to highlight specific characters.

The following abbreviations are used in the species descriptions: A = antennal segments, S = metasomal sterna, and T = metasomal terga. Subgeneric concepts follow Pisanty et al. (2022b) with the necessary modifications detailed below. In diagnoses, the defining characters of a species are given, with those of the indicated comparison species given in parentheses.

Collections studied

AMC Personal collection of Andreas Müller, Wädenswil, Switzerland;

CMHC Carlos M. Herrera collection, Estación Biológica de Doñana, Seville, Spain;

FIOS Personal collection of Francisco Javier Ortiz-Sánchez, El Ejido, Spain;

FLOW FLOWer lab collection, University of Coimbra, Coimbra, Portugal;

EBDC EBD-CSIC collection, Estación Biológica de Doñana, Seville, Spain;

MNHN Muséum pational d'Histoire paturalle, Paris, France.

MNHN Muséum national d'Histoire naturelle, Paris, France;

MRSN Museo Regionale di Scienze Naturali di Torino, Turin, Italy;
 MSC Personal collection of Maximillian Schwarz, Ansfelden, Austria;
 MZUR Zoological Museum of Sapienza University of Rome, Rome, Italy;

NHMUK Natural History Museum, London, United Kingdom;

OÖLM Oberösterreiches Landesmusum, Linz, Austria;

RMNH Naturalis Biodiversity Center, Leiden, the Netherlands;
 SMFD Naturmuseum Senckenberg, Frankfurt am Main, Germany;
 TJWC Personal collection of Thomas J. Wood, Mons, Belgium;

UMONS Laboratory of Zoology collection, University of Mons, Mons, Belgium;

ZISP Zoological Institute of the Russian Academy of Sciences, St. Petersburg, Russia;

ZMHB Museum für Naturkunde, Berlin, Germany.

Results

Genetic results and taxonomic changes

Genetic study of Iberian *Andrena* resulted in a number of important necessary changes to species concepts, as well as further supporting recent decisions. These results are presented here by subgenus; not all Iberian *Andrena* subgenera are treated here, as no taxonomic problems were detected for the majority of species.

Subgenus Aciandrena Warncke, 1968

In Iberia, this is a species-poor subgenus, consisting solely of *A. fulica* Warncke, 1974 and *A. vacella* Warncke, 1975. Warncke described *A. astrella* Warncke, 1975 from Iberia, and used it in combination with *A. fulica* as a subspecies. The two taxa were synonymised by Wood et al. (2020b) as one of the distinguishing characters was that North African populations had males with the clypeus entirely black, and Iberian populations had males with the clypeus yellow-marked, but males with both colour forms can be found in both geographical regions. There are no clear structural differences in the tergal punctation or the structure of the male genital capsule. Genetically (Fig. 1), Iberian and Moroccan populations differ by an average genetic distance of 1.61% (range 1.31–2.09%). Although Iberian material (including specimens WPATW318-21 and WPATW354-21 which are black-faced males from the Sierra Nevada) forms a clade with bootstrap support of 100, the overall low genetic distance is not considered to be sufficient to justify species status. The synonymy of Wood et al. (2020b) is therefore maintained. The broad *A. fulica* clade as a whole is well-defined with bootstrap support of 98.

Subgenus Aenandrena Warncke, 1968

This subgenus was found to be paraphyletic by Pisanty et al. (2022b), and the same result is found here with a COI-based analysis (Fig. 1). *Andrena hystrix* Schmiedeknecht, 1883 falls away from *A. aeneiventris* Morawitz, 1872 and *A. hedikae* Jäger, 1934. For convenience, these taxa are retained in the same subgenus, but future work will be needed to resolve this issue and probably to describe a new subgenus for the species around *A. hystrix* (currently three Palaearctic species from Morocco to Central Asia). Type material for *A. hystrix* was recently rediscovered, and a lectotype for this species is designated below.

In the true *Aenandrena*, four species are currently recognised, of which two are widely distributed, *A.* (*Aenandrena*) *aeneiventris* Morawitz, 1872 that was described from Italy and *A.* (*Aenandrena*) *hedikae* Jäger, 1934 that was described from the western Balkans. Both of the widespread species are nominally distributed from Iberia and Morocco to Central Asia in dry and warm parts of the Palaearctic (Gusenleitner and Schwarz 2002). Barcode analysis (Fig. 1) showed that *A. aeneiventris* specimens from Austria, Hungary, Israel, and Portugal formed a clade with bootstrap support of 99,



Figure 1. Phylogenetic tree (maximum likelihood) of *Andrena* from the subgenera *Aciandrena* Warncke, 1968, *Aenandrena* Warncke, 1968 sensu lato, *Graecandrena* Warncke, 1968, *Poecilandrena* Hedicke, 1933, and the *numida*-group based on the mitochondrial COI gene. *Andrena* (incertae sedis) *relata* Warncke, 1975 is used as an outgroup. Numbers adjacent to branches represent bootstrap support (values of <75 are omitted).

and with low average intraspecific genetic distance of 0.78% (range 0.30–1.06%). In contrast, *A. hedikae* formed two clades. Five specimens from Portugal and Spain formed a clade with bootstrap support of 93 with low differentiation of 0.82% (range 0.16–2.01%). Two specimens from northern Morocco had identical sequences and formed a clade with bootstrap support of 98. These two clades were separated by an average genetic distance of 4.55% (range 4.34–4.72%), and both were well-separated from *A. aeneiventris*, by an average genetic distance of 5.86% (range 4.98–6.67%) for Iberian specimens and 5.20% (range 4.86–5.75%) for Moroccan specimens. These genetic distances are substantial, but no taxonomic action is taken here, as sequences from south-eastern Europe are needed to i) further understand barcode variation in

A. hedikae and ii) identify which of these two clades matches populations from the locus typicus. Moreover, there are no apparent morphological differences between Iberian and Moroccan 'hedikae'. Further study is required.

Subgenus Chlorandrena Pérez, 1890

This subgenus is clearly supported genetically (Pisanty et al. 2022b) and morphologically by the presence of a row of teeth on the posterior face of the female femur combined with strong 'crater punctures' on the terga. Barcodes generally supported all species concepts (Fig. 2), but there are some issues that must be discussed.

Andrena (Chlorandrena) livens Pérez, 1895 was described from north-eastern Spain. Warncke described A. livens algeria Warncke, 1967 from Tunisia on the basis of slight

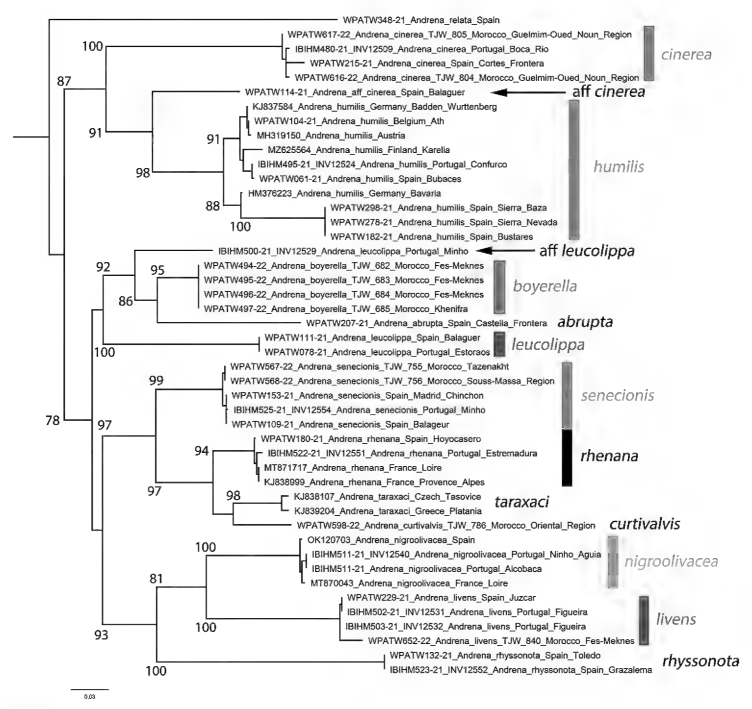


Figure 2. Phylogenetic tree (maximum likelihood) of *Andrena* subgenus *Chlorandrena* Pérez, 1890 based on the mitochondrial COI gene. *Andrena* (*Nobandrena*) *funerea* Warncke, 1975 is used as an outgroup. Numbers adjacent to branches represent bootstrap support (values of <75 are omitted).

differences in the genital capsule and sternum eight. He also described *A. livens gruen-waldti* Warncke, 1967 from Sicily. Based on the sequences here, a Moroccan specimen conforming to *A. livens algeria* shows relatively low differentiation from Portuguese and Spanish specimens, being separated by an average genetic distance of 2.37% (range 2.16–2.47%). Additional samples are required to investigate the status of these subspecies, and also to sample *A. (Chlorandrena) agnata* Warncke, 1967 which is a poorly understood species in an Iberian context; I have seen no Iberian specimens, and it is included on the list on the basis of the single female paratype from Madrid, collected 6.vi.1946 by Dusmet (Warncke 1967). *Andrena agnata* is much closer morphologically to *A. livens* than the nearest neighbour in the current tree, *A. (Chlorandrena) nigroolivacea* Dours, 1873; no taxonomic action is taken until such sequences are available, and a broad *A. livens* concept is maintained for now.

The status of A. (Chlorandrena) boyerella Dours, 1872 and A. (Chlorandrena) leucolippa Pérez, 1895 and the relationship of these taxa to each other has been confused. Warncke (1967) used a subspecies concept, with A. boyerella s. str. present in North Africa and A. boyerella leucolippa present in Iberia and France, A. leucolippa being described from south-western France, locus typicus Riscle (Fig. 3). It is important to note that Pérez writes in his catalogue under entry 442 that he had males from Algeria; in any case, the designation of the lectotype by Warncke (1967) from France fixes this name unambiguously on European populations. North African and Iberian specimens differ, with A. leucolippa presenting denser tergal punctation, in the male sex the yellow colouration of the clypes extends onto the mandibles and the lower paraocular areas (only the clypeus is yellow-marked in A. boyerella), and there are slight differences in the genital capsule. Gusenleitner and Schwarz (2002) listed A. leucolippa as a distinct species.

However, this classification is not immediately stable due to the fact that the type series of A. boyerella is lost, as is the case for all of Dours' types. Dours (1872) described A. boyerella, and there is no doubt over the morphological identity of this taxon based on the clear description. However, Dours gives a distribution of southern France and Algeria. It is therefore the case that, if there are two different species, then Dours' type series was polytypic. Warncke (1967) and Gusenleitner and Schwarz (2002) give the *locus* typicus of A. boyerella as Algeria, but since there is no holotype or designated lectotype, the *locus typicus* is undefined. Genetically, sequences from four North African specimens from the Middle Atlas in Morocco were identical and formed a clade (Fig. 2) that was strongly separated from two sequences from Estorãos near Fafe in northern Portugal [WPATW078-21] and Balaguer in Catalonia in north-eastern Spain [WPATW111-21] by an average genetic distance of 13.27% (range 13.27–13.27%). The two clades are not sisters, being separated by A. (Chlorandrena) abrupta Warncke, 1967. The matter would appear to be clear, that two taxa are present. However, a specimen from Vieira do Minho in northern Portugal [IBIHM500-21] diverged from both these lineages, being separated from the Moroccan sequences by 8.95% and the two other Iberian sequences by 14.51%. This site is only 16 km from the Estorãos site. Morphologically, there are no obvious differences; all three Iberian specimens are males. Because A. leucolippa was described from south-western France, the sequence from north-eastern Spain is here

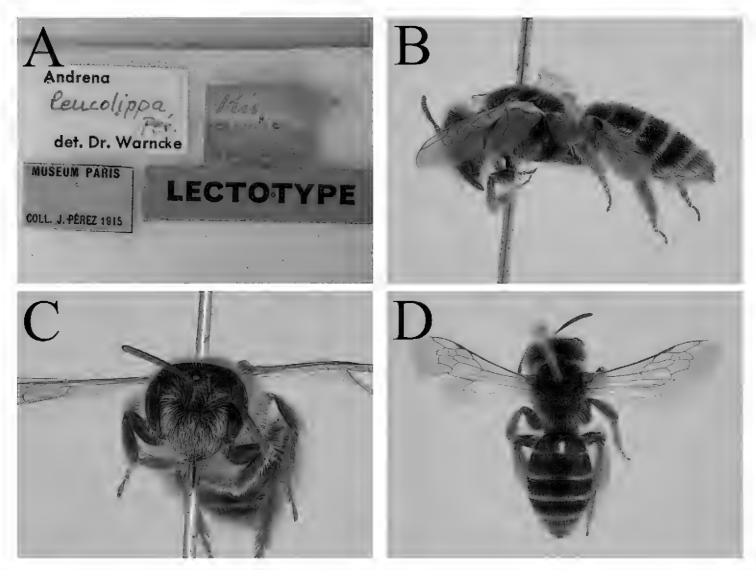


Figure 3. Andrena (Chlorandrena) leucolippa Pérez, 1895 female lectotype **A** label details **B** profile **C** face, frontal view **D** dorsal view.

assumed to be representative of true *A. leucolippa*. For now, the divergent Portuguese sequence is considered to be aberrant until it can be independently replicated. It is possible that this 'aberrant' sequence is a NUMT (nuclear sequences of mitochondrial origin) and represents amplification of nuclear DNA rather than mitochondrial. Since only a single sequence is available, and it is not possible to determine the nature of this sequence, no action is taken. It is not impossible that a third undescribed species is present, but without additional genetic data no further action is taken. Morphology supports the existence of only two species, and in order to fix the name *A. boyerella* on North African populations, a neotype is designated below from barcoded Moroccan material.

Andrena (Chlorandrena) humilis Imhoff, 1832 is the most widespread West Palaearctic Chlorandrena, and it is quite variable over its range. A broad species concept has been used as no consistent morphological differences can be found because of this variability. Sequences from Austria, Belgium, Finland, Germany, Portugal, and Spain formed a broad A. humilis clade (Fig. 2), though intraspecific variation was high at an average of 4.59% (range 0.00–8.33%). Specimens from Iberian mountain chains (Sistema Central, Sierra de Baza, Sierra Nevada) were the most strongly divergent, differing by 8.02–8.33% from specimens from Austria, Belgium, and Germany. However, multiple clades were found without clear geographic structuring. Therefore, a broad species concept is maintained

here, and this overall A. humilis clade has bootstrap support of 98. In Iberia, A. humilis forms a species pair with A. (Chlorandrena) cinerea Brullé, 1832. Andrena cinerea specimens from Morocco, Portugal, and Spain formed a distinct clade with low intraspecific variation of 0.77% (range 0.00–1.54%). This clade was strongly separated from the broad A. humilis clade by an average distance of 15.05% (range 13.89–15.74%). However, a single specimen from Balaguer in north-eastern Spain which was originally identified as A. cinerea diverged from both the broad A. humilis clade by 11.70% (range 11.11–12.65%) and A. cinerea by 12.35% (range 12.35–12.35%). As for the potentially aberrant specimen of A. leucolippa, it is not clear what this sequence represents, though it is not A. (Chlorandrena) kamarti Schmiedeknecht, 1900 from North Africa and Sicily as it also diverges from Moroccan A. kamarti sequences (to be published in the upcoming North African revision). It may also be a NUMT; it is unknown if certain Andrena subgenera are more likely to generate NUMTs, but *Chlorandrena* may potentially represent one such case, with difficult to interpret results generated in the study of the East Mediterranean fauna (G. Pisanty, unpublished data). More study using additional genetic markers is necessary to understand why two seemingly aberrant strong divergent sequences have been generated from Iberian Chlorandrena specimens.

Finally, the *taraxaci*-group (see Schwenninger 2015) formed a monophyletic clade, with *A.* (*Chlorandrena*) *rhenana* Stöckhert, 1930 clearly separated from *A.* (*Chlorandrena*) *taraxaci* Giraud, 1861 which has its western range limit in Central Europe. The taxonomic concepts of Schwenninger (2015) are therefore supported and followed.

Material examined. *Andrena leucolippa*: France: Riscle [43.6564°N, -0.0894°W], 1♀, MNHN (lectotype; Fig. 3).

Subgenus Didonia Gribodo, 1894

Andrena (Didonia) mucida Kriechbaumer, 1873 is a highly unusual species. It is bivoltine, with the first generation seemingly specialising on Muscari (Asparagaceae) and the second generation specialising on genera from the former Dipsacaceae (now Caprifoliaceae) such as Scabiosa. This specialisation is associated with a morphological change, with females of the first generation possessing tibial scopae composed of simple hairs, and females of the second generation possessing tibial scopae composed of plumose hairs. There is the possibility that these generations may actually represent distinct species, as for putatively bivoltine taxa like A. (Holandrena) decipiens Schenck, 1861 that was found to consist of two taxa (Mandery et al. 2008). However, barcodes from females from the first and second generations from central and southern Spain show that this is not the case, with an average intraspecific genetic distance of 0.30% (range 0.15–0.46%; Fig. 4). The spring and summer generation individuals from Guadalajara were separated by 0.15%, strongly supporting the position that A. mucida is a bivoltine species displaying the currently unique trait of intergenerational variation in the structure of the pollen collecting hairs.

Additionally, A. (Euandrena) solenopalpa Benoist, 1945 was previously placed in the subgenus Didonia (Warncke 1968a). Genetic evidence instead places this taxon in the subgenus Euandrena (Bossert et al. 2022). This placement is a better fit, as

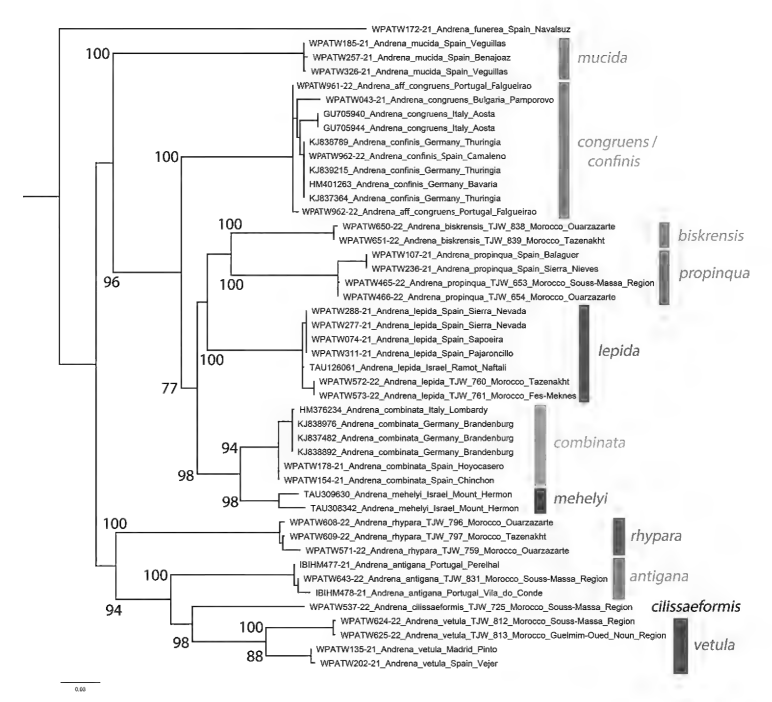


Figure 4. Phylogenetic tree (maximum likelihood) of *Andrena* from the subgenera *Didonia* Gribodo, 1894 and *Simandrena* Pérez, 1890 based on the mitochondrial COI gene. *Andrena* (*Nobandrena*) *funerea* Warncke, 1975 is used as an outgroup. Numbers adjacent to branches represent bootstrap support (values of <75 are omitted).

A. solenopalpa has typical Euandrena foveae (narrow and narrowing further ventrally), the male A3 slightly exceeds the length of A4+5, and the process of the labrum is not so strongly thickened and elongate as in A. mucida; this simply represents convergence on a similar morphology. It is possible that the subgenus Didonia is monotypic, containing only A. mucida, but this requires additional study.

Material examined. *Andrena mucida*: SPAIN: Guadalajara, Veguillas, 2 km N, Barranco de la Isa, 17.v.2021, 1♀, leg. T.J. Wood, TJWC [BOLD accession number WPATW185-21], on *Muscari* spp.; Málaga, Benaoján, Cueva del Hundidero, 3.vi.2021, 1♀, leg. T.J. Wood, TJWC [BOLD accession number WPATW257-21], on *Scabiosa atropurpurea*; Guadalajara, Veguillas, CM-1006, 9.vii.2021, 1♀, leg. T.J. Wood, TJWC [BOLD accession number WPATW326-21], on *Scabiosa atropurpurea*.

Subgenus Euandrena Hedicke, 1933

In comparison to the situation in the Eastern Mediterranean (Praz et al. 2019), Iberian *Euandrena* are comparatively well-resolved, with lower species diversity, fewer undescribed species, and clearer species boundaries. However, unrecognised montane species are present (Wood et al. 2021), and there are some problems related to the group of species lumped together under a broad *A. bicolor* Fabricius, 1775. This current work does not deal with the complex issue concerning the two clades of *A. bicolor* as identified by Praz et al. (2019), but instead with the status of material from southern Spain and North Africa. Material identified as *A. bicolor* s.l. from the Col du Zad in the Middle Atlas of Morocco at an altitude of 2100 m [WPATW387-22] and the southern slopes of the Sierra Nevada above Trevélez in Spain at an altitude of approximately 1800 m [WPATW368-21] fell close to *A. angustior* (Kirby, 1802) and far from *A. bicolor* s.l. (Fig. 5). The specimen from the Sierra Nevada was caught on the same *Campanula* (Campanulaceae) plant as specimen WPATW290-21 which falls into the *A. bicolor* s.l. clade. These two Spanish specimens are separated by a genetic distance of 9.80%, and clearly cannot be conspecific despite their strong morphological similarity.

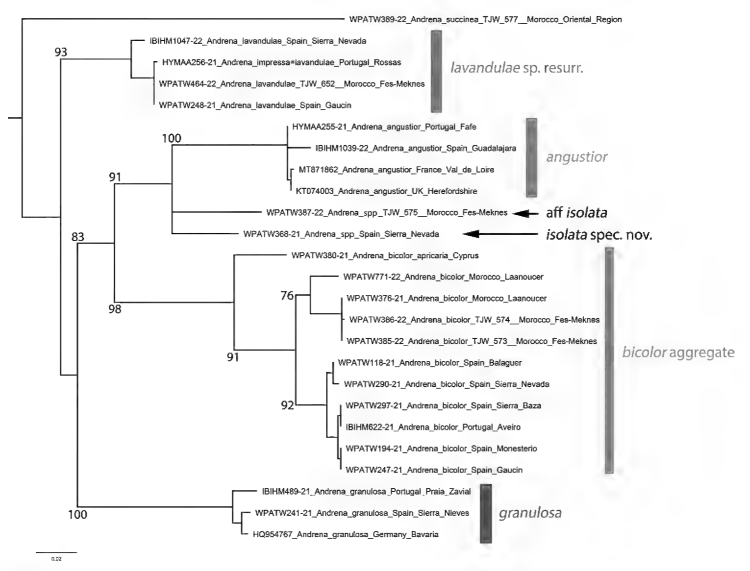


Figure 5. Phylogenetic tree (maximum likelihood) of *Andrena* subgenus *Euandrena* Hedicke, 1933 based on the mitochondrial COI gene. *Andrena* (*Pruinosandrena*) *succinea* Dours, 1872 is used as an outgroup. Numbers adjacent to branches represent bootstrap support (values of <75 are omitted).

More broadly, the Sierra Nevada specimen was separated from the *A. bicolor* s.l. clade by an average genetic distance of 9.32% (range 7.95–9.80%) and the Moroccan specimen was separated from the *A. bicolor* s.l. clade by an average distance of 9.06% (range 8.50–9.80%). These two specimens were separated from each other by a distance of 4.99%. The species from the Sierra Nevada is described as new to science below, but the name to apply to the North African taxon (present also in Tunisia based on barcode data, Praz, in litt.) is complex and requires discussion. The nomenclatural and taxonomic status of *A. bicolor* s.l. will be resolved in a future revisionary work dedicated to the subgenus *Euandrena* at a West Palaearctic scale.

Andrena (Euandrena) bicolor and associated Euandrena species in North Africa

Euandrena specimens with black and orange pubescence can be found infrequently across north-western Africa. They have typically been referred to as A. bicolor in the literature, but two distinct taxa are present. One corresponds to A. bicolor s.l., but the other is clearly distinct genetically; morphologically it can be distinguished by the structure of the clypeus, with A. bicolor s.l. with the clypeus shiny between the punctures, whereas in the second taxon the puncture interspaces are shagreened and dull, and there is a weak longitudinal furrow, similar to what can be seen in species like A. (Euandrena) angustior, though more apically situated and occupying a shorter distance.

Warncke (1974) used subspecific concepts for dealing with A. bicolor, with two subspecies used for North African material - A. b. oraniensis Lepeletier, 1841 and A. b. agraria Warncke, 1974 (see also distribution maps in Gusenleitner and Schwarz 2002). However, the use of these names is not immediately straightforward. Andrena oraniensis Lepeletier, 1841 was described from Oran in northern Algeria (Lepeletier 1841) and is a confused and unclear taxon. In the MNHN collection, the type series cannot be located, and it may have been lost as many specimens have been moved between different boxes by past workers and curators. In the Pérez collection, there are long series of Andrena (Melandrena) florentina Magretti, 1883 that were identified by Pérez as A. oraniensis. Warncke (1967) noted this, but argued that this must be incorrect due to size, and instead placed the name in combination with A. bicolor as a subspecies found in North Africa. Warncke (1974) later authored the replacement name A. bicolor agraria Warncke, 1974 for Andrena (Euandrena) nigriventris Pérez, 1902 which was described from northern Algeria and Morocco (locus typicus Tangier, following lectotype designation) but which is a preoccupied name, nec. Apis nigriventris Gmelin, 1790 which is a synonym of A. (Melandrena) nitida (Müller, 1776). Warncke then used both oraniensis and agraria as subspecies, giving an overlapping distribution in Warncke (1974, e.g. both occurring at Oukaimeden in the High Atlas), but in his distribution maps (see Gusenleitner and Schwarz 2002) giving a non-overlapping distribution, with oraniensis more or less occurring only in Algeria and Tunisia and agraria occurring only in Morocco.

In the original description of *A. oraniensis*, Lepeletier (1841: 245) draws attention to the colour of the hairs on the hind legs. Specifically, he states that: "cuisses des deux postérieures garnies des poils ferrugineux pales; leurs jambes et leurs tarses à poils noirs en

dessus, ferrugineux en dessous". This bicoloured tibial scopa (dark dorsally, ferruginous ventrally) does not correspond at all to members of the A. bicolor s.l. group which have uniformly orange tibial scopae. It does however correspond very well to the concept of A. florentina which has a distinctive bicoloured scopa, one of the characters that allows it separation from its sister taxon A. bicolorata (Rossi, 1790). Based on the identifications made by Pérez, who probably saw the original specimen in Lepeletier's collection and Lepeletier's original description, the position is taken that A. oraniensis cannot be a Euandrena taxon displaying the colour pattern of A. bicolor s.l. However, without a type, it is undesirable to make A. oraniensis the senior synonym for A. florentina. Andrena oraniensis is therefore declared a nomen dubium until such time as the original syntypic series can be located.

The next oldest available name from North Africa is A. bicolor agraria. Pérez (1902, as A. nigriventris) writes: "Abdomen très luisant, particulièrement les depressions, que précèdent les bourrelets très prononcés". This is suggestive, as the distinct North African taxon has strongly depressed tergal margins. However, barcoded A. bicolor s.l. from Morocco also show this character, so it is not diagnostic in and of itself. Examination of the lectotype specimen of A. nigriventris (designated by Warncke 1967; Fig. 6) shows that the clypeus is apically smooth and shing between the punctures (Fig. 6C), meaning that it cannot belong to the distinct taxon, and it remains a synonym of A. bicolor

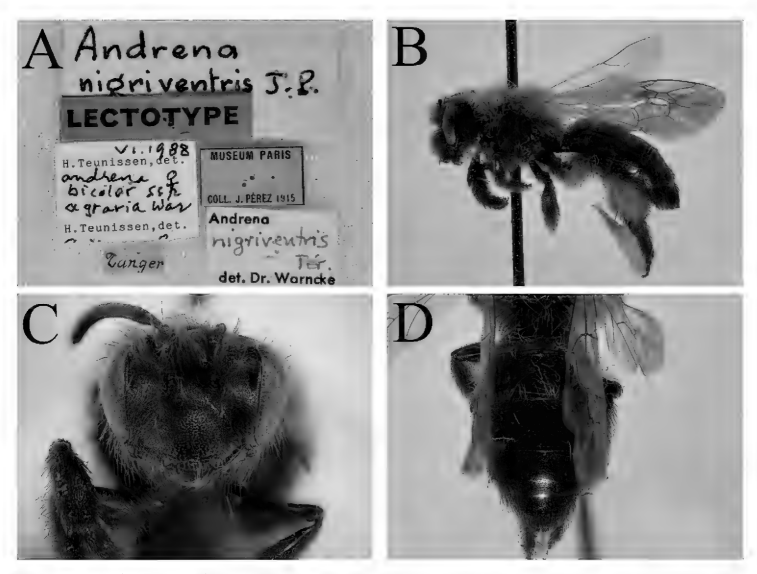


Figure 6. Andrena (Euandrena) nigriventris Pérez, 1902 (nec. Gmelin, = Andrena bicolor agraria Warncke, 1974) female lectotype **A** label details **B** profile **C** face, frontal view **D** terga, dorsal view.

s.l. One additional name is available, that of *A.* (*Euandrena*) *fervida* Pérez, 1902, a taxon described from Algeria. The lectotype is in poor condition, lacking its metasoma (Fig. 7). Diagnosis is therefore challenging, but the clypeus is shiny between the punctures and it cannot be conspecific with the distinct North African taxon. The position of Warncke (1967) that this name is synonymous with *A. bicolor* s.l. is maintained. The distinct North African taxon would therefore appear to be undescribed. No taxonomic action is taken in the current work – further genetic data will either demonstrate a closer affinity with the specimen from the Sierra Nevada, in which case this name will be available to apply to North African populations, or these sequences will confirm its distinct nature, and it can be described. In any case, a name is required for the Iberian lineage which is unambiguously undescribed.

Material examined. *Andrena bicolor*: ALGERIA: Theniet El Had [35.8727°N, 2.0007°E] (1♀, MNHN (lectotype of *A. fervida*); MOROCCO: Tanger [35.7537°N, -5.7906°W], 1♀, MNHN (lectotype of *A. nigriventris* Pérez).

Andrena (Euandrena) lavandulae Pérez, 1902, sp. resurr.

Andrena (Euandrena) lavandulae Pérez, 1902: 156 \$\rightarrow\$ [France, lectotype by present designation: MNHN].

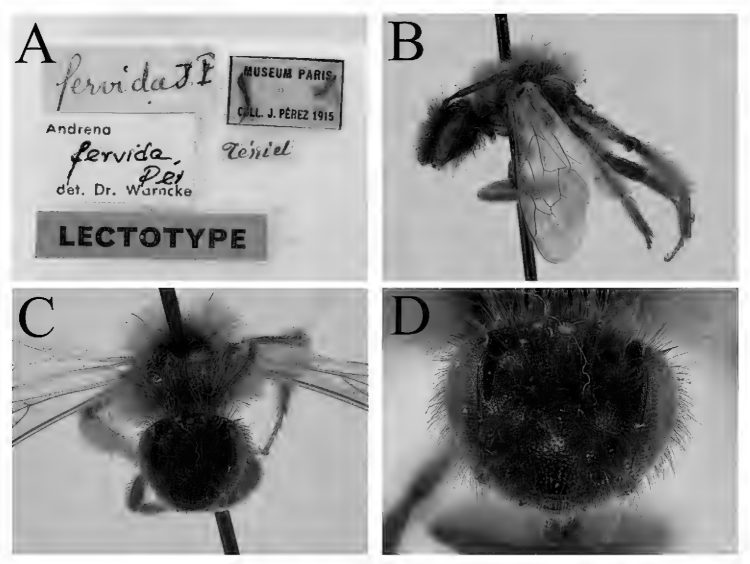


Figure 7. Andrena (Euandrena) fervida Pérez, 1902, female lectotype **A** label detail **B** profile **C** face, frontal view **D** face detail, frontal view.

Andrena (Euandrena) angustior impressa Warncke, 1967: 234, ♀♂ [Morocco: OÖLM, examined] syn. nov.

Remarks. Warncke (1967) synonymised *A. lavandulae* with *A. bicolor* without apparently inspecting the type. This is not explicitly clear in the text of Warncke (1967), but in this publication he typically reported lectotypes in the Paris collection when he designated them, and usually labelled paratypes as well, often taking a selection for his personal collection. There are no specimens of *A. lavandulae* in the Warncke collection, which combined with the lack of mention of a lectotype in Warncke (1967) leads me to believe that he did not inspect the type series. He then described *Andrena angustior impressa* Warncke, 1967 in the same publication, a taxon that was subsequently elevated to species status by Wood et al. (2021) based on genetic and morphological evidence.

In the MNHN collection, specimens of *A. lavandulae* are labelled as lectotype and paralectotype by Teunissen (August 1986; Fig. 8), but these designations were never published. The female specimen is therefore recognised here as a lectotype, by present designation. Examination of these specimens shows that they are clearly the same taxon as *A. impressa* and distinct from *A. bicolor*, with the weakly depressed, shagreened, and weakly shing tergal margins, the black terminal fringe and hairs flanking the pygidial plate, the intermixed light and dark hairs on the face, and in the male sex by the broadened gena (broader than the width of the compound eye). The lectotype comes from Banyuls-sur-Mer in the south-west of France, which falls within the distribution of the taxon as presented by Wood et al. (2021). Additional genetic sequences from Spain and Morocco closely match the Portuguese sequence of *A. impressa* presented by Wood et al. (2021), forming a well-supported clade (Fig. 5; bootstrap support of 93) that is distinct from both *A. angustior* and *A. bicolor. Andrena impressa* syn. nov. is therefore synonymised with *A. lavandulae* sp. resurr. The distribution is Morocco, Algeria, Portugal, Spain, and France (Wood et al. 2021).

Material examined. FRANCE: Banyuls [Banyuls-sur-Mer, 42.5658°N, 2.8658°E], 1♂, 1♀, MNHN (female lectotype, by present designation).

Subgenus Graecandrena Warncke, 1968

This subgenus is also species-poor in Iberia, containing only *A. impunctata* Pérez, 1895, *A. montarca* Warncke, 1975, *A. nebularia* Warncke, 1975, and *A. verticalis* Pérez, 1895. Two taxa are uncommonly collected in Iberia (*A. impunctata* and *A. montarca*). *Andrena nebularia* was considered to be endemic to Spain, but new collections in Morocco have demonstrated its presence in a small part of the Middle Atlas. Genetically, there is almost no differentiation, with the Moroccan specimens separated by 0.26% and 0.52% (Fig. 1).

Material examined. *Andrena nebularia*: Morocco: Fès-Meknès, Boulemane, 5 km SE, junction of R503 and N4, 1900 m, 19.v.2022, 2♀, leg. T.J. Wood, TJWC; Fès-Meknès, Boulemane, R503, 7 km SE of Boulemane, 1900 m, 1♂, 6♀, 22.v.2022, leg. T.J. Wood, TJWC; Fès-Meknès, Boulemane, R503, SE of Ait Karmosse, 1750 m, 22.v.2022, 1♂, leg. T.J. Wood, TJWC.

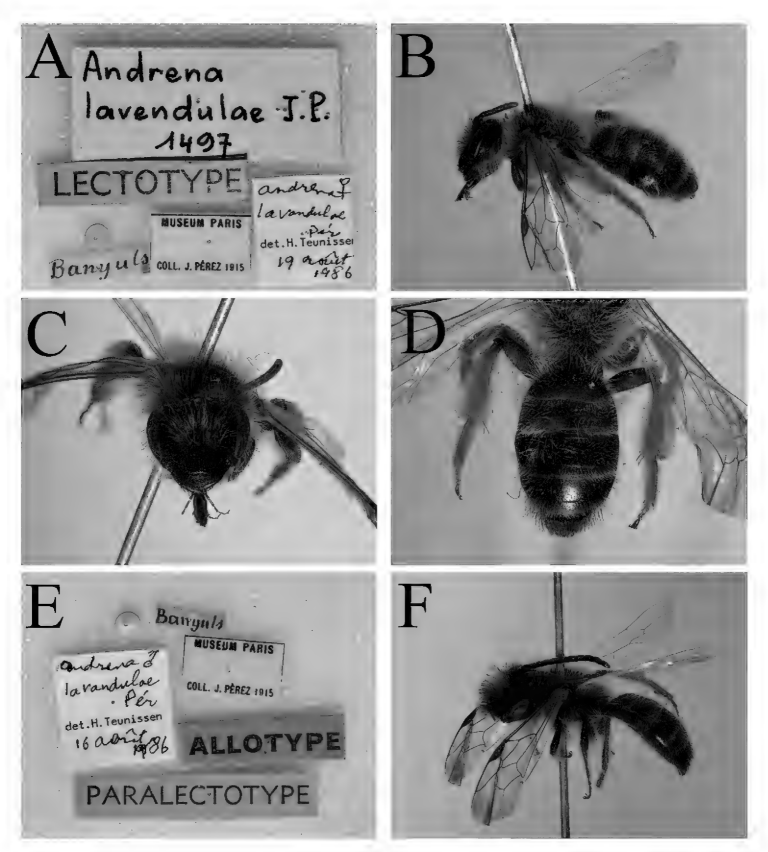


Figure 8. Andrena (Euandrena) lavandulae Pérez, 1902, female lectotype A label details B profile C face, frontal view D terga, dorsal view; male syntype E label details F profile.

Subgenus Melanapis Cameron, 1902

Andrena (Melanapis) fuscosa Erichson, 1835

Andrena fuscosa was described from southern Spain by Erichson (1835). Andrena fuscosa has had a complicated taxonomic history, and has been described many times due to its variation in colouration and size (Gusenleitner and Schwarz 2002). Iberian genetic data are therefore of interest since the peninsula represents the *locus typicus*.

Iberian sequences were identical with each other, and closely matched sequences from Israel and India (the latter identified as A. (Plastandrena) agilissima Scopoli, 1770 but clearly misidentified; Melanapis was actually described as a genus based on material from India), being separated by an average of 1.53% (range 0.88–1.82%; Fig. 9). They were more clearly separated from Moroccan sequences, by an average of 5.43% (range 3.94–5.93%). Given the lack of morphological differences between North African and Iberian specimens, a broad interpretation of this species is taken here. Including all sequences together, average intraspecific distance is 3.14% (range 0.00–6.23%) which is considered to be acceptably small. Additional study is necessary to establish whether North African material is consistently distinct; the first nominally available name would be A. (Melanapis) rutila Spinola, 1838 which was described from Egypt.

Subgenus Melandrena Pérez, 1890

This subgenus is strongly derived within *Andrena*, but it shows substantial morphological variation that has led to the description of the subgenera *Hyperandrena* Pittioni, 1948 and *Zonandrena* Hedicke, 1933, both of which are now placed within an expanded *Melandrena*. There are several problems within this subgenus in an Iberian context.

Andrena (Melandrena) morio Brullé, 1832

Andrena (Melandrena) morio Brullé, 1832: 353, ♀♂ [Greece: MNHN, not examined]. Andrena (Melandrena) hispania Warncke, 1967: 212, ♀♂ [Spain: OÖLM, examined] syn. nov.

Remarks. Andrena hispania Warncke, 1967 was described from Algeciras in southern Spain. The identification characters given by Warncke are comparatively weak and rely on hair colour and the degree of infuscation of the wings, without mentioning definitive structural characters. Genetically, the three *A. hispania* sequences from Spain and Portugal mixed with A. morio sequences from Israel, Morocco, Portugal, Spain, Tunisia, and Turkey without forming a cluster, this group having bootstrap support of 100 (Fig. 10). Confusingly, additional sequences from Greece, Morocco, and Tunisia formed two sister groups to this clade. There is no clear geographic pattern, and so a broad A. morio concept is adopted, including A. hispania syn. nov. as a direct synonym of A. morio. Additional genetic data using more powerful techniques are required before an alternative taxonomic conclusion can be drawn. Andrena morio is known to be highly variable in its colouration, leading to a higher than average number of synonyms for an Andrena species (Gusenleitner and Schwarz 2002). Andrena hispania was classically thought of to have only a single generation in the spring, which may explain why its wings are less infuscate than A. morio, particularly in the summer generation of this species that can be noticeably darker than spring flying individuals.

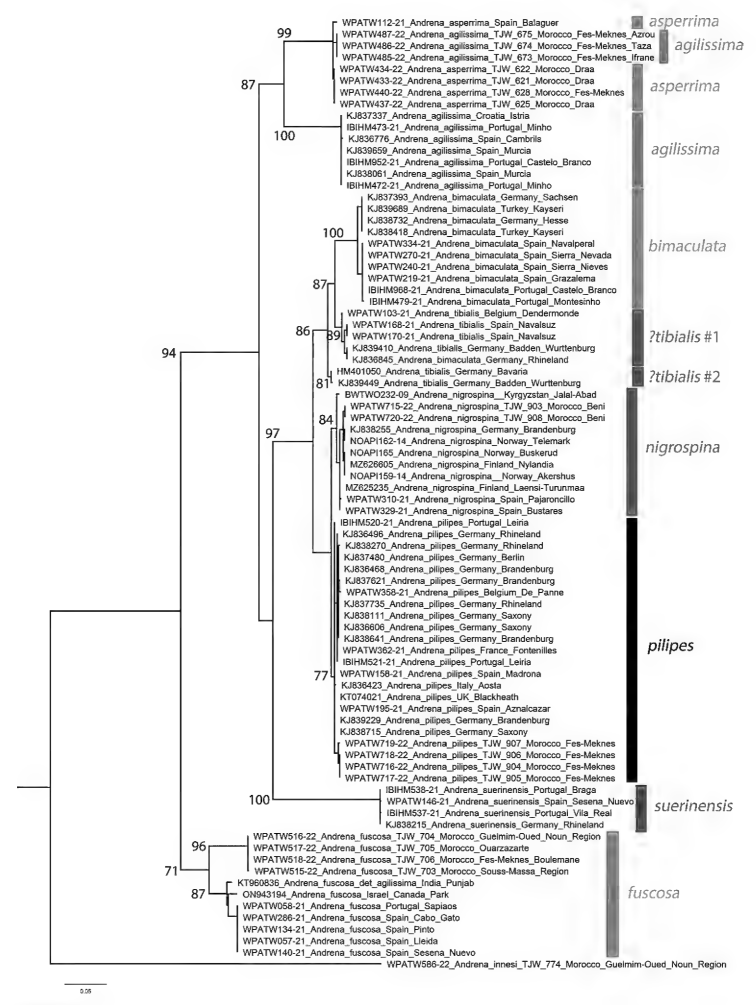


Figure 9. Phylogenetic tree (maximum likelihood) of *Andrena* from the subgenera *Melanapis* Cameron, 1902, *Plastandrena* Hedicke, 1933, and *Suandrena* Warncke, 1968 based on the mitochondrial COI gene. *Andrena* (incertae sedis) *innesi* Gribodo, 1894 is used as an outgroup. Numbers adjacent to branches represent bootstrap support (values of <75 are omitted).

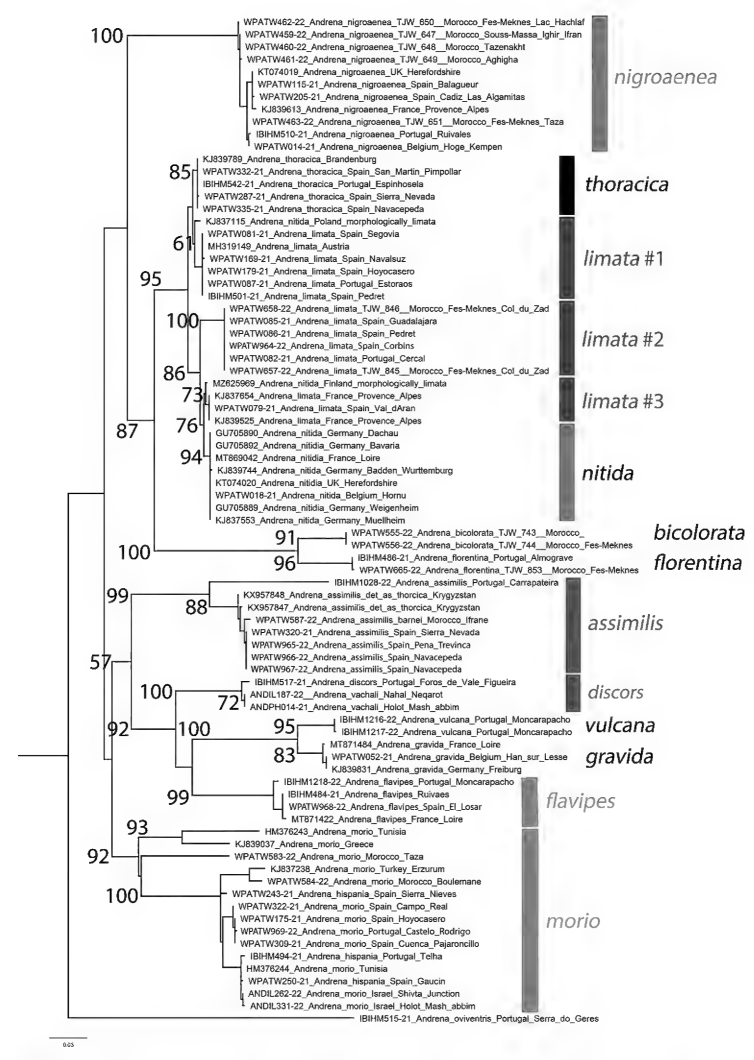


Figure 10. Phylogenetic tree (maximum likelihood) of *Andrena* subgenus *Melandrena* Pérez, 1890 based on the mitochondrial COI gene. *Andrena* (*Ovandrena*) *oviventris* Pérez, 1895 is used as an outgroup. Numbers adjacent to branches represent bootstrap support (values of <75 are omitted).

Andrena (Melandrena) nitida (Müller, 1776), Andrena (Melandrena) thoracica (Fabricius, 1775), and Andrena (Melandrena) limata Smith, 1853

Remarks. The specific differences between these taxa is unclear across southern Europe. In some northern countries, only two taxa are present (A. nitida and A. thoracica, e.g. the United Kingdom), with no introgression observed. Andrena nitida flies only in the spring, whereas A. thoracica is bivoltine, flying in both the spring and the summer. In Central Europe, A. limata can be found, this taxon also being bivoltine. Differentiation between the three taxa in Central Europe has often utilised hair colouration characters, as in this region the three taxa are separable with reference to the hairs on the sides of the mesosoma (light in A. nitida and A. limata, dark in A. thoracica) and the hairs of the face and the tibial scopal (light and dark in A. nitida, uniformly dark in A. limata and A. thoracica). There are additional characters such as the colour of the hind tibial spur and the density of the punctures T1, but these are less commonly referred to; both A. nitida and A. limata have dense punctures on T1 (separated by up to 1 puncture diameter), whereas they are more clearly spaced in A. thoracica (punctures separated by 1–2 puncture diameters). It is important to note that A. limata is a replacement name for A. lucida Lepeletier, 1841 nec. A. lucida Panzer, 1798 which is nominally a synonym of A. bicolor Fabricius, 1775 but this must be established (see section on subgenus Euandrena). Andrena lucida Lepeletier was described from 'France', without further details. I have not been able to inspect the type which should be in the MNHN collection, but which I could not locate.

The situation in southern Europe is much more challenging. In south-western Europe, *A. limata* becomes much darker, and therefore closely resembles the colour form of *A. thoracica*, with extensive black pubescence on the mesosoma laterally. This colour form was described as *A. limata mixtura* Warncke, 1967 (illustrated by Wood et al. 2020a). This form can be recognised by the dense punctures of T1, separating it from Iberia *A. thoracica*; the colour of the hind tibial spur is variable and of little consistent utility. Because of this variation, Warncke later adopted a much broader concept of *A. nitida*, with *A. nitida* s. str., *A. nitida limata*, and *A. nitida mixtura* (e.g. Warncke 1974). This position was not followed by Gusenleitner and Schwarz (2002) who followed a three distinct species interpretation (*nitida, limata* including *mixtura, thoracica*).

Analysis of barcodes does not provide clarity (Fig. 10). Whilst *A. thoracica* and *A. nitida* form reciprocally monophyletic clades, three different clades are formed for *A. limata*. There is no geographic pattern to these clades; clade #1 contains individuals from Austria, Poland (KJ837115; specimen identified as *A. nitida* but associated photograph on BOLD shows *A. limata* colour pattern), Spain, and Portugal; clade #2 contains individuals from Morocco, Spain, and Portugal; and clade #3 contains individuals from Finland (MZ625969; specimen identified as *A. nitida* but associated photograph on BOLD shows *A. limata* colour pattern), France, and Spain. Confusingly, sequences IBIHM501-21 and WPATW086-21 come from two different female specimens collected from the same field near Pedret i Marzà in north-eastern Spain. As Iberian individuals fall into three of these clades and the power of COI analysis is

clearly insufficient to deal with potential hybridisation and introgression, no further action can be taken on the basis of these results. A dedicated study using more powerful genetic markers is necessary to resolve this issue. For now, I maintain the position of Gusenleitner and Schwarz (2002) in that there are three species in this group (*nitida*, *limata* including *mixtura*, and *thoracica*). It will probably be necessary for a future study to designate a neotype for *A. limata* if Lepeletier's original specimens cannot be definitively located.

Andrena (Melandrena) discors Erichson, 1841

Andrena (Melandrena) discors Erichson, 1841: 192, ♀ [Algeria: ZMHB, examined]. Andrena (Melandrena) creberrima Pérez, 1895: 46, ♀♂ [Algeria: MNHN, examined] syn. nov.

Andrena (Melandrena) vachali Pérez, 1895:

[Tunisia, lectotype by present designation: MNHN] syn. nov.

Andrena (*Melandrena*) *bodemeyeri* Benoist, 1969: 246, ♀ [Tunisia: MNHN, examined].

Remarks. There has been extensive confusion between A. discors and A. creberrima. Erichson (1841) described A. discors from Algeria; a specimen is preserved in the ZMHB collection which may be automatically the holotype, but this is ambiguous as it is unclear if Erichson described the species from multiple specimens (Fig. 11). In any case, the specimen is a syntype, and Erichson described the species only in the female sex. The specimen conforms to the classical concept of the species, having the typical features of the former members of the subgenus Zonandrena, i.e. a clypeus with punctures forming a pattern of weak longitudinal channels. The pubescence of the face is black (Fig. 11C), and there are numerous intermixed dark hairs on the scutum (Fig. 11D). The tibial scopae are orange (Fig. 11E), and there are weak indications of light tergal hair bands but these are degraded and scarcely visible (Fig. 11F). Warncke (1974) gave numerous localities for A. discors from Morocco, Algeria, and Tunisia. He listed A. aff discors from Madrid based on two specimens (Warncke 1976), but later discarded these records, giving only two points in southern Spain in his distribution maps in addition to the points from north-western Africa (Gusenleitner and Schwarz 2002). I have examined 14 female and six male specimens of A. discors, all from southern parts of Portugal and Spain (Algarve, Alto Alentejo, Cádiz, Huelva, Málaga, Sevilla), where it is active from late February to April.

The situation is confused due to the status of two additional taxa described by Pérez, A. (Melandrena) creberrima Pérez, 1895 and A. (Melandrena) vachali Pérez, 1895. Andrena creberrima was described from Algeria like A. discors. Warncke (1967) designated a lectotype for A. creberrima (Fig. 12), but did not understand the concept of this species, listing only the type locality of Bône in Warncke (1974). He later gave a distribution map of Algeria (the holotype point), Italy, and Greece (Crete). Examination of the lectotype shows that the specimen is slightly paler than the type of A. discors, with fewer dark intermixed hairs on the scutum (Fig. 12B), and the tergal hair bands

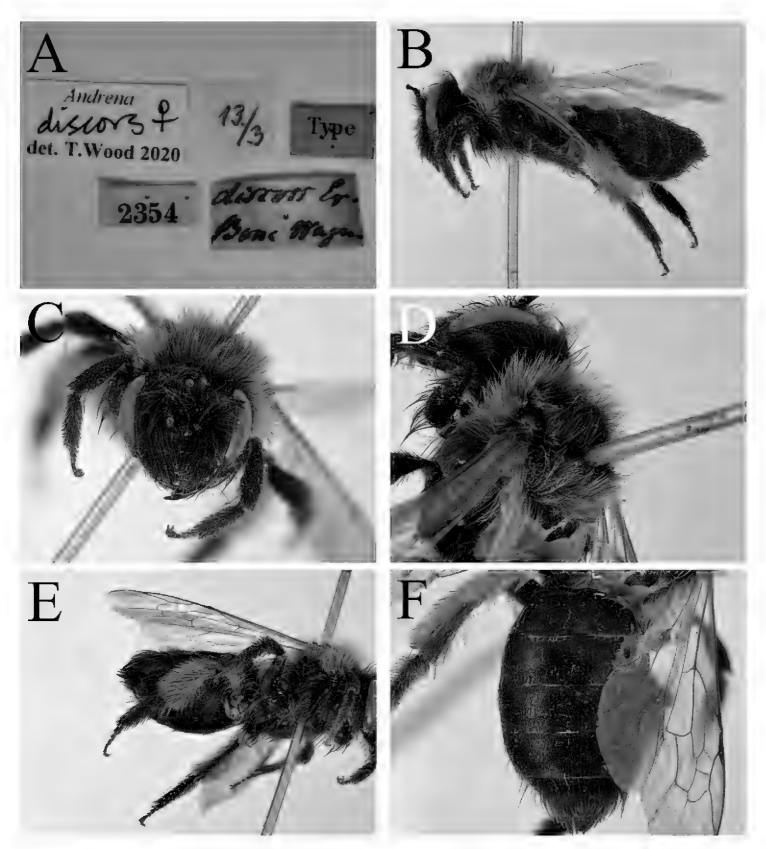


Figure 11. Andrena (Melandrena) discors Erichson, 1841, female holotype/syntype **A** label details **B** profile **C** face, frontal view **D** scutum, dorsolateral view **E** tibial scopa, profile view **F** terga, dorsal view.

in better condition and therefore slightly more pronounced (Fig. 12D). This can be contrasted with the type of *A.* (*Melandrena*) bodemeyeri (Fig. 13; known synonym of *A. discors*, see Warncke 1967) which is comparatively dark. This kind of variation in hair colour is typical for the subgenus *Melandrena*, and has led to the large number of synonymous names being described, for example for *A.* (*Melandrena*) flavipes Panzer, 1799.

Grouping A. creberrima under A. discors is straightforward. The problem comes with A. vachali which was classically considered to be distinct from A. discors and with a greater range extending from the Canary Islands to southern Israel (Gusenleitner

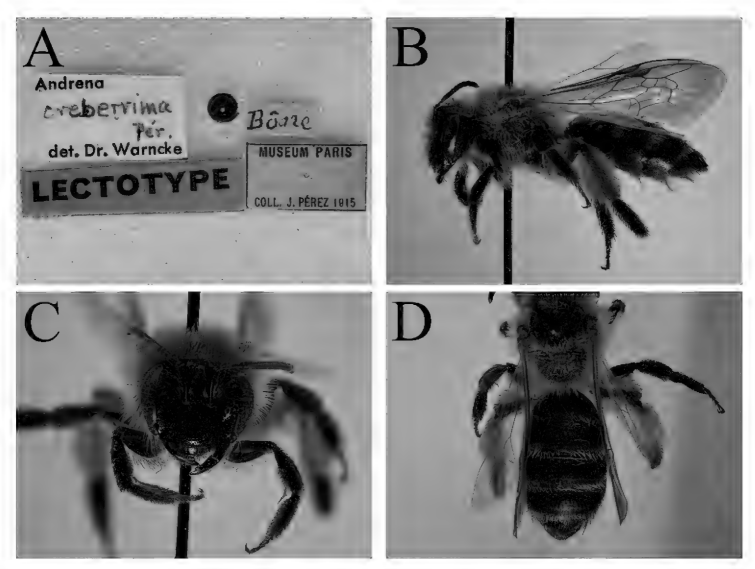


Figure 12. Andrena (Melandrena) creberrima Pérez, 1895, female lectotype **A** label details **B** profile **C** face, frontal view **D** terga, dorsal view.

and Schwarz 2002). Moreover, type material for A. vachali has not actually been designated, as Warncke (1967) could not find material in the MNHN collection. Examination of the MNHN showed several specimens of A. vachali in the Vachal collection from Foum Tatahouine [Tataouine, Tunisia] labelled "Andrena vachali type Pérez" in the handwriting of Vachal (Fig. 14). One specimen was labelled as a lectotype with a handwritten 'A. vachali' determination label by H. Teunissen (Fig. 14A), but this designation was never published. Pérez (1895) did not mention the collection locality, and Warncke (1967) indicated that the species was described from Algeria. However, Gusenleitner and Schwarz (2002) give the *locus typicus* as Tunisia. Examination of the catalogue of Pérez did not reveal an entry for A. vachali, as is often the case, either because Pérez did not write one, or because the information is hidden under a name that differs from the published name (see below for the case of A. (Plastandrena) asperrima Pérez, 1895). It is therefore very difficult to know the type locality with certainty. However, on the basis that the MNHN specimens are labelled as 'type' by Vachal, they were present in his collection, and the species was named after Vachal, the position is taken that these represent genuine syntypes. The female specimen labelled as 'type' by Vachal is chosen as a lectotype, by present designation. Morphologically, it corresponds to the concept of A. vachali, with black pubescence on the face and mesepis-

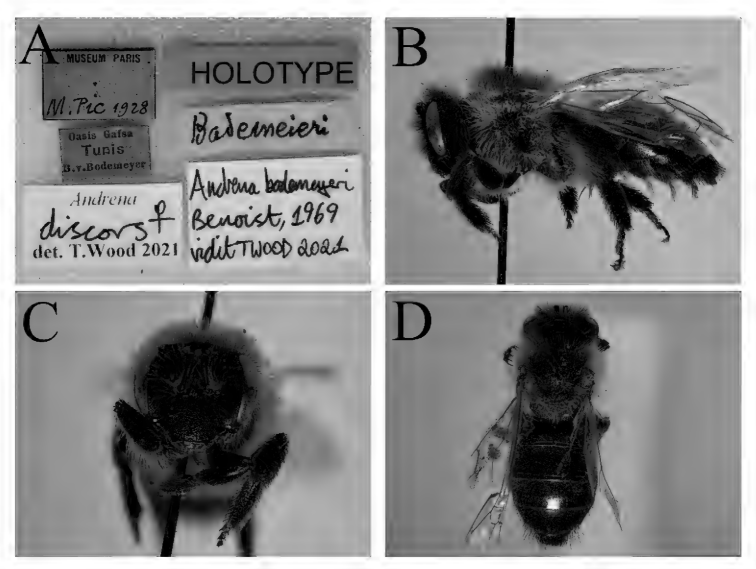


Figure 13. Andrena (Melandrena) bodenmeyeri Benoist, 1969, female holotype **A** label details **B** profile **C** face, frontal view **D** dorsal view.

ternum, with reddish-brown pubescence on the remaining parts of the mesosoma, and with predominantly reddish-brown tibial scopae, though this specimen shows some darker hairs dorsally (c.f. Fig. 13D); specimens of *A. vachali* can be found with entirely orange-red tibial scopae. Comparison of sequences from *A. discors* from southern Portugal and two *A. vachali* from southern Israel show a very small difference of 0.91% (range 0.91–0.91%; Fig. 10). These three specimens group together and have posterior support of 100, and are clearly separated from the three other former *Zonandrena* species that group together in this clade, namely *A. flavipes*, *A. (Melandrena) gravida* Imhoff, 1832, and *A. (Melandrena) vulcana* Dours, 1873.

Taking a step back, the differentiation between *A. creberrima*, *A. discors*, and *A. vachali* has almost always been based on colouration, with *A. discors* the darkest, *A. vachali* the lightest, and *A. creberrima* somewhat intermediate. Males are generally rarer in collections; taken together, I have examined 215 female specimens, but only 73 male specimens of this *discors-creberrima-vachali* group. However, examination of males has led me to the conclusion that there are no apparent differences in the males of *A. creberrima*, *A. discors*, and *A. vachali*, and indeed the male of only one of these nominal taxa was actually originally described. All have white hairs over the majority of the face with clear black hairs laterally along the inner margins of the compound eyes,



Figure 14. *Andrena (Melandrena) vachali* Pérez, 1895, female lectotype **A** label details **B** profile **C** face, frontal view **D** terga, dorsal view.

a genital capsule that is typical of the former *Zonandrena* with the dorsal surface of the gonocoxae with granular shagreen, with the penis valves moderately broad, and without an emargination in the outer margins of the gonostyli (Fig. 15). Indeed, the genital capsule of a syntype of *A. creberrima* (Fig. 15B), the barcoded *A. discors* from southern Portugal [IBIHM517-21] (Fig. 15C), and *A. vachali* specimens from Morocco have identical genital capsules (Fig. 15D). Moreover, numerous 'aberrant' female individuals can be found across this range with more or less pronounced tergal hair bands and variably dark or light pubescence, particularly in Crete where individuals corresponding in different ways to the concepts of *A. creberrima*, *A. discors*, and *A. vachali* can be found.

Based on this colour variation, the lack of variation in the male genital capsule, the unclear and overlapping distributions given by Warncke (*A. creberrima* and *A. vachali* both reported from Crete, *A. creberrima* and *A. discors* both reported from Algeria), and the very low genetic distance between Portuguese and Israeli specimens, both *A. creberrima* syn. nov. and *A. vachali* syn. nov. are synonymised with *A. discors*. This resolves this long-running confusion as to the correct name and identity of this taxon (Baldock et al. 2018; Wood et al. 2020a).

Distribution. Portugal, Spain (Canary Islands, mainland), Morocco, Algeria, Tunisia, Italy, Malta, Libya, Greece (Crete), Egypt, Israel, Jordan.

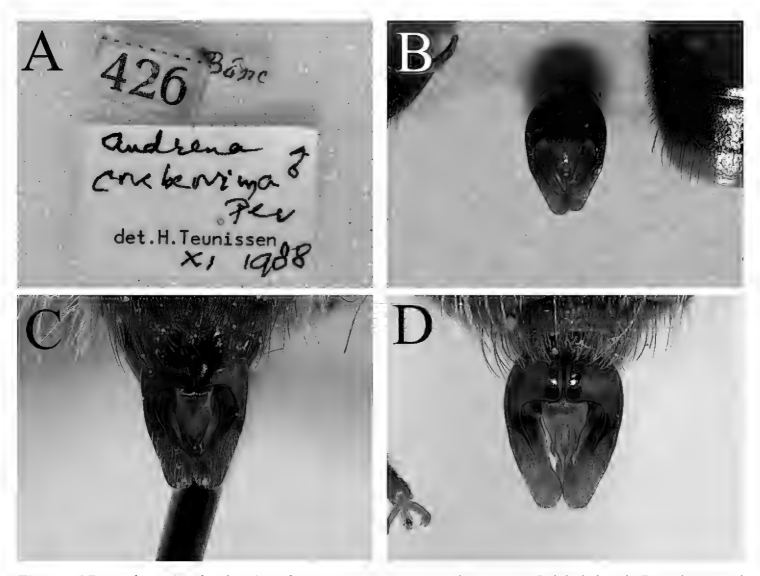


Figure 15. Andrena (Melandrena) creberrima Pérez, 1895, male syntype **A** label details **B** male genital capsule; Andrena (Melandrena) discors Erichson, 1841 **C** male genital capsule; Andrena (Melandrena) vachali Pérez, 1895 **D** male genital capsule.

Material examined (illustrative). ALGERIA: label information illegible, possibly 'Bone' [= Annaba], 1♀, ZMHB (holotype/syntype of A. discors); Bône [=Annaba, 36.9092°N, 7.7264°E], 1♀, MNHN (lectotype of A. creberrima); Alger, Birmandreis [Bir Mourad Raïs], 24.iii.-30.iv.1910, 13, 12, leg. J. Bequaert, ZMHB; GREECE: Krete, Anidri b. Paleochora, 200 m, 31.iii.2002, 12, leg. A. Müller, AMC; Krete, Palaeochora, 0–50 m, 29.iii.2002, 52, leg. A. Müller, AMC; Kreta, Kavalos [Kavallos], 21.v.1986, 12, leg. Brande, OÖLM; Crete, Dept. Heraklion, Moulia, S. of Aghia Varvara, 26.iv.1973, 12, leg. v. Ooststroom, RMNH; ITALY: Lazio, Roma, Via Falcognana, 8.iv.1990, 1♀, leg. G.G.M. Schulten, RMNH; Roma, Via Falcognana, 15.iv.1993, 1♀, leg. G.G.M. Schulten, RMNH; ISRAEL: Negev, 15 km S of Be'er Sheva, 31.iii.1988, 2, leg. R. Leys, RMNH; **Jordan:** Petra, 800 m, 28.ii.-4.iii.1986, $1 \circlearrowleft$, $2 \circlearrowleft$, leg. K.M. Guichard, NHMUK; **Malta:** Buskett [woodlands, \exists ad-Dingli], 7.iii.2018, 1, M. Balzan Colln.; Morocco: Drâa-Tafilalet, Tazenakht, Anezal, 5 km NE Ait Igga, 21.iv.2022, 13, 42, leg. T.J. Wood, TJWC; Souss-Massa, R105, Tizirt, 8 km N, Ighir Ifran env., 12.iii.2022, 1♂, 1♀, leg. T.J. Wood, TJWC; PORTUGAL: Foros de Vale Figueira, Montemor-o-Novo, 20.iii.2019, 1&, leg. T.J. Wood, TJWC; Montemor-o-novo, Foros de Vale de Figuera, 24.iv.2011, 1♀, leg. A. Albernaz-Valente,

TJWC; **SPAIN:** Canary Islands, Lanzarote, Haria, 6.ii.1979, 1\$\(\frac{1}{3}\), leg. Ellis & Simon-Thomas, RMNH; Bollullos Par del Condado (Huelva), 160 m, 28.iii.2009, 1\$\(\frac{1}{3}\), leg. F.J. Ortiz-Sánchez, FJOS; Puerto de Gáliz, P.N. Alcornocales (Cádiz), 425 m, 28.iv.2007, 1\$\(\frac{1}{3}\), leg. F.J. Ortiz-Sánchez, FJOS; Cádiz, Zahara de los Atunes, 12.iii.1977, 1\$\(\frac{1}{3}\), leg. Madey, NHMUK; **Tunisia:** Foum Tatahouine [32.9361°N, 10.4458°E], 1\$\(\frac{1}{3}\), MNHN (lectotype of *A. vachali*, by present designation); Oasis Gafsa [34.4350°N, 8.7678°E], 1928, 1\$\(\frac{1}{3}\), leg. B. v. Bodemeyer, MNHN (holotype of *A. bodemeyeri*); Fritissa, 17–19. iv.1983, 1\$\(\frac{1}{3}\), TJWC.

Subgenus Micrandrena Ashmead, 1899

This is by far the most species-rich subgenus in Iberia, containing 37 species. Pisanty et al. (2022b) took the decision to synonymise the subgenera *Distandrena* Warncke, 1968, *Fumandrena* Warncke, 1975, and *Proxiandrena* Schmid-Egger, 2005 with a broad concept of *Micrandrena*, as morphological differences such as the shape of the foveae, propodeal triangle, and presence or absence of striations on the clypeus are inconsistent and insufficient to allow unambiguous differentiation between the groups, as well as lacking phylogenetic support. This broad *Micrandrena* definition is followed here.

Andrena (Micrandrena) obsoleta Pérez, 1895 and Andrena (Micrandrena) mariana Warncke, 1968 sensu lato

The treatment of *A. obsoleta* has a long and confused history that has caused many problems. Warncke (1967) designated a female lectotype from Babor in northern Algeria (Fig. 16). However, this lectotype designation is incorrect. In the catalogue of Pérez, under entry 1506 (p. 214), Pérez writes: "Sicile, \bigcirc - M'pellier \bigcirc . - Tebessa, avril, \bigcirc . -Biskra, mars, $3 \updownarrow$, rares. – Bône, \updownarrow . – Marseille, \circlearrowleft . - Bône, $27/\text{iv} \circlearrowleft \updownarrow$ in cop. Souci et autres [illegible – presumably a type of flower]". Babor is not mentioned, and so Warncke's lectotype designation is rejected. Examination of specimens labelled as obsoleta by Pérez from the localities indicated shows that they belong to a different taxon to that of Warncke's false lectotype. Warncke's false lectotype is a female *Distandrena* (=*Micran*drena) with a clearly striate clypeus, and T1 is polished. This conforms to the concept of A. (Micrandrena) nitidula Pérez, 1903, and indeed Warncke (1967) synonymised A. nitidula with A. obsoleta. Only female specimens of A. obsoleta from Bône [=Annaba] in northern Algeria could be found in the Pérez collection, and almost all were in very poor condition, missing their metasomas. A single female from Bône was however in good condition, and it is here designated as a new lectotype (Fig. 17). This new lectotype specimen (i.e. the true A. obsoleta) morphologically falls within the A. (Micrandrena) mariana Warncke, 1968 complex within the former Distandrena, as it lacks striations on the clypeus. The larger A. (Micrandrena) merimna Saunders, 1908 can be excluded as this species has a very long A3 that clearly exceeds the length of A4+5, whereas members of the A. mariana complex have A3 at most slightly exceeding A4+5, A4 and A5 are sub-quadratic and slightly shorter than wide, A6-12 are as long as wide.

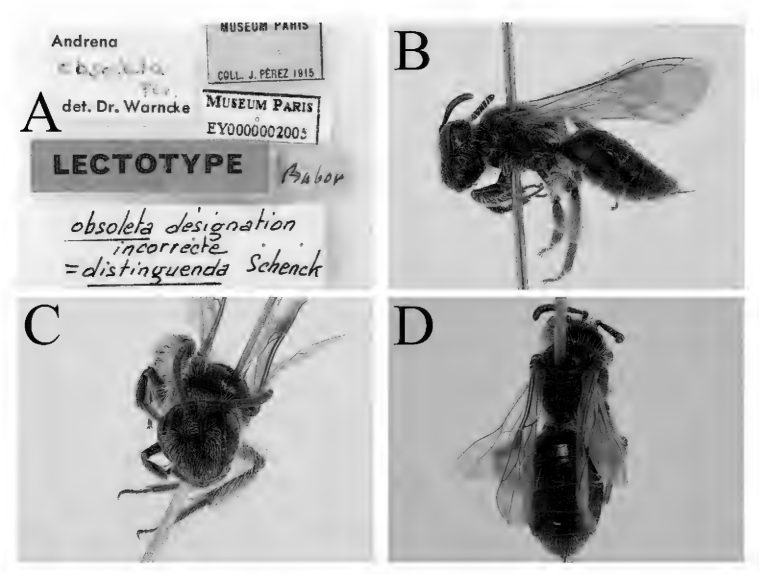


Figure 16. Andrena (Micrandrena) obsoleta Pérez, 1895, false female lectotype, =Andrena (Micrandrena) nitidula Pérez, 1903 A label details B profile C face, frontal view D dorsal view.

Warncke described A. mariana s. str. from the island of Fuerteventura in the Canary Islands, stating that the species could potentially be found in Morocco (Warncke 1968b). He then described five subspecies in subsequent publications (Warncke 1974, 1975a): A. m. mica Warncke, 1974, A. m. leptura Warncke, 1974, A. m. solda Warncke, 1974, A. m. alma Warncke, 1975, and A. m. tenostra Warncke, 1975. For North African taxa, Warncke (1974) described A. mariana solda from Morocco (Tangier) and Algeria (Algiers, including Husseyn-Dey [Hussein Dey] the *locus typicus*, Constantine, Bone, Blida) and Tunisia (Carthago), and noted that this species corresponds to the Andrena forms from Bone identified as iota, kappa, and lambda by Saunders (1908), though this is clearly wrong for *lambda* as the genital capsule illustrated by Saunders shows A. (Micrandrena) djelfensis Pérez, 1895. The A. mariana solda taxon differs from A. mariana s. str. in the female sex by the darker tergal margins, the darker antennae, and the more extensively shagreened clypeus. Warncke (1974) described A. mariana mica from southern Algeria (locus typicus Ghardaia) and Morocco, giving it a more arid distribution. It also differs from A. mariana s. str. in the female sex by the darker tergal margins and darker antennae, but the clypeus is more extensively shiny than in A. mariana solda, and the nervulus is said to be more antefurcal. Andrena m. leptura is a more eastern taxon, being described from Libya and Egypt, and having a broader process of the labrum than any of these species.

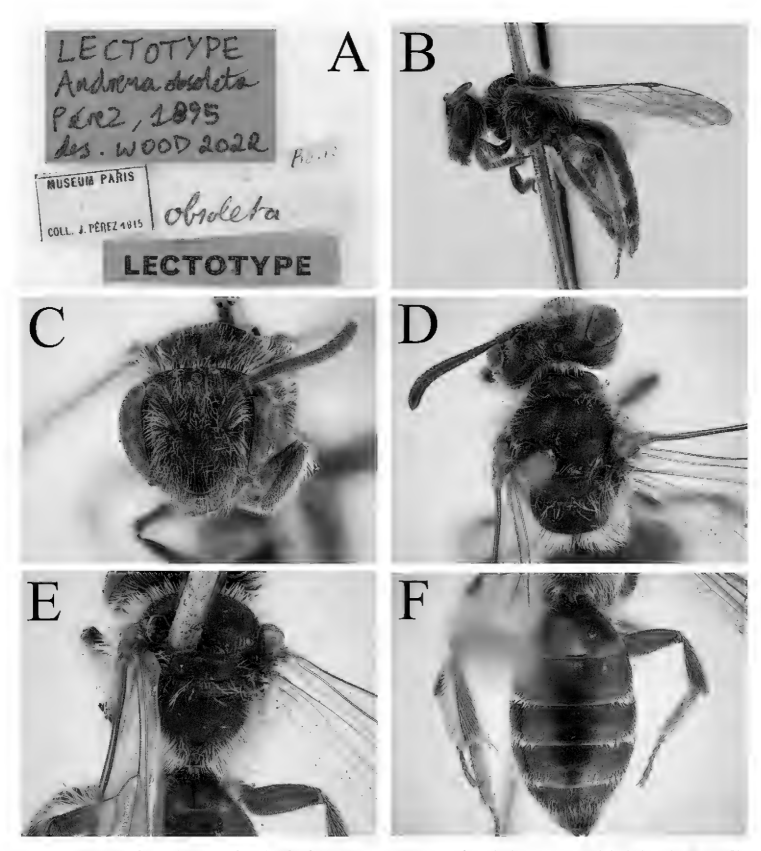


Figure 17. Andrena (Micrandrena) obsoleta Pérez, 1895, true female lectotype **A** label details **B** profile **C** face, frontal view **D** scutum, dorsal view **E** propodeal triangle, dorsal view **F** terga, dorsal view.

Examination of *A. mariana solda* material reveals that it is conspecific with the newly designated lectotype of *A. obsoleta* and is synonymised with it syn. nov. As identified by Warncke, the taxon has a wide distribution across Mediterranean parts of Morocco, Algeria, and Tunisia (Warncke 1974, as *A. m. solda*). Pérez's reference to specimens from France indicates that his original syntypic series would have been polytypic, or he may have changed his mind between writing this catalogue entry and his 1895 publication, or even after this point. Warncke (1974: 40) indicated that *A. m. solda* is present in Sicily which is supported by a newly examined specimen (see below),

though he did not designate any Italian specimens as paratypes. As no specimens from France labelled as *obsoleta* could be found in Pérez collection, the conservative position is taken here that this species does not occur in France. The implications for Warncke's misinterpretation of *A. obsoleta* are discussed below.

As A. mariana solda is actually a synonym of a valid species that was described prior to A. (Micrandrena) mariana s. str., this has implications for the species-status of the other taxa lumped under A. mariana by Warncke. Genetic sequences were available for A. mariana mica Warncke, 1974 from southern Morocco, A. mariana s. str. from southwestern Morocco, and A. mariana alma Warncke, 1975 from Portugal (locus typicus near Córdoba, southern Spain; Fig. 18). Contrary to the position of Warncke (1968b), examination of material from Morocco, Algeria, and Tunisia shows that A. mariana is actually widely distributed across this region, predominantly in southern, more arid areas. Genetically, A. mariana s. str. was more closely related to A. (Micrandrena) abjecta Pérez, 1895, separated by an average genetic distance of 5.55% (range 5.49-5.61%), and was strongly separated from A. m. mica by 12.73% (range 12.50-12.95%) and A. m. alma by 10.94% (range 10.65-11.39%). Andrena abjecta + A. mariana s. str. therefore formed a clade with bootstrap support of 99. Andrena m. mica and A. m. alma formed a clade, but were strongly separated from each other by 11.26% (range 10.97-11.41%). The conclusion therefore is that each of these taxa is distinct, and so A. mica stat. nov. and A. alma stat. nov. are raised to species status. Morphologically, all three taxa can clearly be separated by the structure of the female clypeus, the colour of the antennae, the extent of lightening on the tergal margins, the strength of the scutal punctures, and the shape of the process of the labrum.

Warncke described two further subspecies of *A. mariana*: *A. mariana leptura* from Egypt and *A. mariana tenostra* from south-eastern Spain, the latter specifically from a single female specimen from Villajoyosa in the province of Alicante, then listing additional specimens from Almería and Murcia (Warncke 1976). The status of *A. mariana leptura* will be dealt with in a future publication, as its exact species concept and its eastern and western limits are unclear, though it will be a valid species as it clearly differs from *A. mariana* s. str.; its relationship with *A. mariana* s.l. specimens from southern Israel must also be clarified (Pisanty, in litt.). In Iberia, *A. m. tenostra* is very poorly known. The distinguishing features given by Warncke (1975a) are that the process of the labrum is narrowly trapezoidal and that the clypeus is arched and somewhat flattened in the middle, in contrast to *A. alma* where the clypeus is largely flattened. Additional female specimens were found in Granada which highlight a difference not noted by Warncke, which is that the scutellum is polished and shiny, whereas it is dull in *A. alma*, and more broadly the scutal punctures are less strongly pronounced. Given that subtle differences within this group represent large genetic differentiation, *A. tenostra* stat. nov. is raised to species status.

Finally, a number of specimens were found in the very south of Spain (Málaga, Sevilla) which show a morphology very close to that of *A. mica*. However, ecologically this does not make sense as *A. mica* is not known from the more humid and Mediterranean areas north of the Atlas Mountains. Examination of these specimens shows that A3 is much longer than A4+5, whereas it is as long as A4+5 in *A. mica*. These specimens probably represent an additional undescribed species in the *A. mariana* complex, but



Figure 18. Phylogenetic tree (maximum likelihood) of *Andrena* subgenus *Micrandrena* Ashmead, 1899 based on the mitochondrial COI gene. *Andrena* (incertae sedis) *murana* Warncke, 1975 is used as an outgroup. Numbers adjacent to branches represent bootstrap support (values of <75 are omitted).

they are not described at this time, as it would be beneficial to have barcoded specimens to confirm this differentiation. They are therefore referred to as aff *mica* in the identification key.

It is important to note that this material does not belong to *A. abjecta*, the status of which in Iberia is unclear. Warncke (1976) did not record this species from Iberia, and only his distribution maps (Gusenleitner and Schwarz 2002) indicate the presence of this species in northern and eastern Spain. This distribution does not make sense given that all other known records of *A. abjecta* come from north-western Africa (Morocco, Algeria, Tunisia), so the species would naturally be expected to occur in southern Iberia. Dardón (2014) listed two female specimens from Huelva (22.iv.1987) and Zaragoza (9.iii.1991). The specimen from Huelva may represent the undescribed aff *mica* taxon, but it is more difficult to conclude on the specimen from Zaragoza. For now, *A. abjecta* is retained on the Iberian list, but I have not examined any specimens and I consider its presence in Iberia doubtful.

Material examined. Andrena alma: PORTUGAL: Albandeira, near Lagoa, 20.iv.2005, 1♀, leg. D.W. Baldock, TJWC; Algarve, Casaqueimada (7 km N of Silves), 20.iii.1995, 1♀, leg. T. & M. Simon Thomas, RMNH; Algarve, Monte Gordo, Retur, Praia do Cabeço, 29.iii.2022, 1♂, leg. T.J. Wood, TJWC; Algarve, Tavira, Cacela Velha, 28.iii.2022, 1♂, leg. T.J. Wood, TJWC; Castelo Branco, Fundão Vale de Prazeres, 17.iv.2021, 1♀, leg. C. Siopa, FLOW; SPAIN: El Soldado, Córdoba [38.3747°N, -5.0001°E], 1♀, leg. Seyrig, OÖLM (holotype); Almodóvar del Campo (Ciudad Real), 700 m, 24.iii.2005, 1♂, leg. F.J. Ortiz-Sánchez, FJOS; Santa Ana la Real, Sierra Aracena (Huelva), 630 m, 13.iv.2006, 2♂, leg. F.J. Ortiz-Sánchez, FJOS; El Hongo (P.N. Doñana), 30.iii.2018, 1♂, leg. F. Molina, EBDC.

Andrena mariana s. str.: Algeria: Saida, 15 km S of Sfissifa, Ben Ikhou, st. 6, 6.iv.1983, 6, leg. R. Leys & P. v. d. Hurk, RMNH; 23 km NE of Ras El Ma, st. 38, 26.iv.1983, 1, leg. R. Leys & P. v. d. Hurk, RMNH; Dayet el Kerch [Daïet el Kerch], st. 5, 5.iv.1983, 2, leg. R. Leys & P. v. d. Hurk, RMNH; Morocco: Guelmim-Oued Noun, Guelmim, Asrir, 1 km W, 20.iii.2022, 1, leg. T.J. Wood, TJWC; Spain: Fuerteventura, Valley Granadillos [28.3864°N, -14.0865°W], 17.iv.1934, 1, OÖLM (holotype); Tunisia: Ksar Hadada, 4–5.iv.1996, 46, leg. K. Deneš, OÖLM; M'saken, 20–21.iv.1998, 2, leg. K. Deneš, OÖLM; Wadi Ram, 45 km E Douz, 4.iv.2006, 1, leg. J. Straka, OÖLM.

Andrena mica: Algeria: Ghardaia [32.5047°N, 3.6419°E], 1♀, OÖLM (holotype); Morocco: 30 km E Midelt, 13.v.1995, 1♀, leg. Mi. Halada, OÖLM; Drâa-Tafilalet, Ouarzazate, P1506, Telouet, Adaha, 1700 m, 18.iv.2022, 6♀, leg. T.J. Wood, TJWC; Drâa-Tafilalet, Ouarzazate, 2 km W Agouim, 1800 m, 13.iv.2022, 3♀, leg. T.J. Wood, TJWC; Ifkern, 25 km E Boulemane, 24.v.1995, 1♀, leg. Mi. Halada, OÖLM; M'rirt (30 km N), 11.iii.1989, 1♀, leg. H. Teunissen, RMNH; Tunisia: Hammamet env, 15.iii.1996, 1♀, leg. K. Deneš, OÖLM; Kasserine, 13.iv.1998, 1♀, leg. K. Deneš, OÖLM; Nefta [Naftah], 20.v.1993, 1♀, leg. J. Batelka, OÖLM.

Andrena aff mica: Spain: 40 km W Málaga, Yunquera, 800 m, 29.iv.2003, 5\(\sigma\), leg. J. Halada, OÖLM/TJWC; La Corchuela (Dos Hermanas, Sevilla), 35 m,

27.iii.2009, 2Å, 2♀, leg. F.J. Ortiz-Sánchez, FJOS; Río Blanco, Aguadulce (Sevilla), 300 m, 17.v.2008, 1♀, leg. F.J. Ortiz-Sánchez, FJOS.

Andrena nitidula: Algeria: Babor, 1♀, MNHN (false lectotype of A. obsoleta).

Andrena obsoleta: Algeria: Bône [=Annaba, 36.9092°N, 7.7264°E], 1♀, MNHN (lectotype of A. obsoleta, by present designation); 23 km NE of Ras El Ma, st. 38, 26.iv.1983, 1♀, leg. R. Leys & P. v.d. Hurk, TJWC; 5 km N of Mecheria, Djebel Antar, st. 10, 7.iv.1983, 1♀, leg. R. Leys & P. v.d. Hurk, RMNH; ITALY: Sicilia orient., Taormina, 200 m, Ghardino [Giardini Naxos], 25.iii.1950, 1♂, Hartig & Grisham, MZUR; Tunisia: Tunis, 1911, 3♀, leg. O. Schmiedeknecht, SMFD.

Andrena tenostra: Spain: Villajoyosa [38.5097°N, -0.2299°E], 11.v.1936, 1♀, leg. Andréu, OÖLM (holotype); Salobreña, Granada, 8.v.1983, 3♀, leg. W. Perrandin, OÖLM/TJWC.

Andrena (Micrandrena) distinguenda Schenck, 1871 and Andrena (Micrandrena) nitidula Pérez, 1903

The status of these two species has been extensively argued over, and despite much attention the position remains somewhat unclear. Warncke (1967) argued that A. distinguenda Schenck, 1871 was an unavailable name because it was preoccupied by A. distinguenda Schenck, 1853. He then authored the replacement name A. obsoleta spongiosa Warncke, 1967, using the incorrect concept that A. obsoleta was the oldest available name for this group of species, as discussed above. Schönitzer et al. (1992) clarified the situation, and argued that A. distinguenda Schenck, 1853 was a nomen nudum because it appears without a description in a discussion concerning species related to A. bicolor. As a nomen nudum, the name is available and cannot preoccupy its later usage. Therefore, the name A. distinguenda Schenck, 1871 is both available and valid as the senior synonym; A. obsoleta spongiosa is consequently an unnecessary replacement name.

Andrena distinguenda was described from western Germany, and A. nitidula was described from south-western France (lectotype from Bordeaux [though this is not indicated on the specimen, it bears the number '675' which refers to the entry for A. nitidula in the catalogue of Pérez], designated by Warncke 1967, Fig. 19). As discussed above, Warncke (1967) incorrectly recognised a specimen that morphologically corresponds to A. nitidula as the lectotype of A. obsoleta, and then synonymised A. nitidula with A. obsoleta. He then treated A. distinguenda Schenck, 1871 as a subspecies of A. obsoleta in the form A. obsoleta spongiosa. He then later adopted a three subspecies model, A. o. obsoleta (North Africa), A. o. nitidula (Western Europe), and A. o. spongiosa (Central Europe) (see Schönitzer et al. 1992), finally introducing a fourth subspecies A. o. puella Alfken, 1938 in Sardinia and Sicily (see Gusenleitner and Schwarz 2002). Schönitzer et al. (1992) argue that both A. distinguenda and A. nitidula can be morphologically separated in north-western Italy (Bordighera). They argue that A. nitidula occurs in North Africa, citing Pérez (1903) and Alfken (1914). They also argue that specimens from southern Spain (Tarifa) appear to be somewhat intermedi-

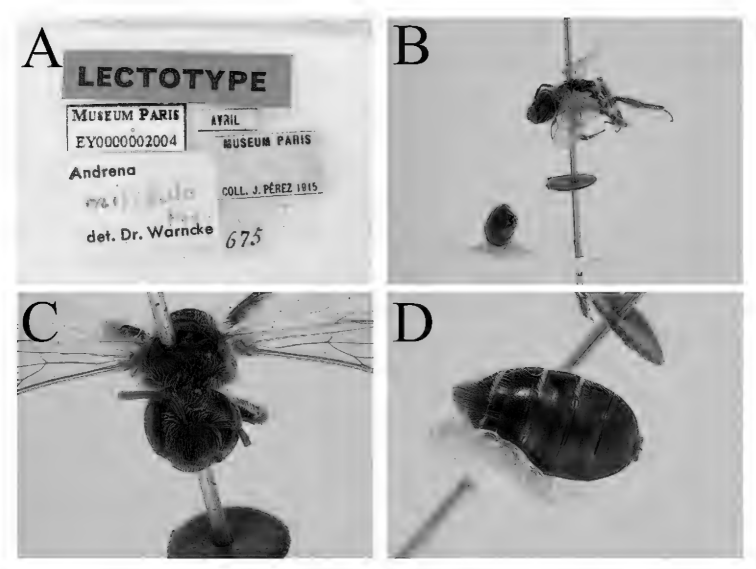


Figure 19. Andrena (Micrandrena) nitidula Pérez, 1903, female lectotype A label details B profile C face, frontal view D terga, dorsal view.

ate between A. distinguenda and A. nitidula, concluding that the question as to whether there are two species or simply a very variable single species remains open.

Burger and Herrmann (2003) revised European material from this species pair using the names A. distinguenda and A. nitidula, after having examined syntypes of A. nitidula in the ZMHB collection. Using morphological characters, they conclude that (in a European context) A. distinguenda has a predominantly eastern distribution, from Central Europe to Greece, with populations also in the south of France and a few records from north-eastern Spain. In contrast, A. nitidula is a western species, from Iberia to the south and east of France and into south-western Germany and the north-west of Italy. They reach the same conclusion as Schönitzer et al. (1992), that where these two forms occur in sympatry (Germany, France, Italy) they can be morphologically separated. They do not conclude on the status of *A. obsoleta*, but note that A. nitidula occurs in North Africa, and that Warncke's treatment is unclear. This problem arises because it does not appear that any of the authors of Schönitzer et al. (1992) or Burger and Herrmann (2003) inspected the lectotype of A. obsoleta designated by Warncke in the MNHN. This lectotype bears a handwritten label that the lectotype designation is incorrect, and that the specimen corresponds to A. distinguenda Schenck (Fig. 16A). However, it is unclear who wrote this, as neither Schönitzer et al. (1992)

nor Burger and Herrmann (2003) mention this explicitly, and I do not recognise the handwriting on the label.

Barcode analysis complicates this matter further (Fig. 18). Analysis of sequences from France, Germany, Morocco, Portugal, and Spain shows the presence of two clear clades. One clade is composed of specimens identified as *A. nitidula* from France, Portugal, and Spain, including specimens from the south of Iberia (Algarve, Málaga, Murcia). This clade would appear to be the true *A. nitidula*. However, the second clade contained sequences from specimens identified as *A. distinguenda* from central Germany, but also from a specimen identified as *A. nitidula* from central Spain (Guadalajara) and two specimens identified as *A. nitidula* from northern Morocco (Fès-Meknes, Oriental). These two clades were consistently separated genetically, by an average genetic distance of 7.08% (range 5.58–8.77%). The nominally true *A. nitidula* clade showed low intraspecific variation of 0.26% (range 0.00–0.63%). However, the situation in the clade containing *A. distinguenda* sequences from Germany showed intraspecific variation of 2.84% (range 0.00–4.10%), with the 4.10% genetic distance recorded between specimens from Germany and the Middle Atlas (Bakrit, near Azrou) in Morocco.

Morphologically, both Spanish and Moroccan specimens falling into the *A. distinguenda* clade conform to the concept of *A. nitidula* using the criteria specified by Burger and Herrmann (2003). Indeed, all material I have identified from northwestern Africa morphologically conforms to the concept of *A. nitidula* sensu Burger and Herrmann. Although the genetic data strongly supports the existence of two species, the conflict between the genetic and morphological results suggests that, at the moment, they cannot be consistently separated morphologically across their range. No taxonomic action is taken here, and the morphological criteria of Burger and Herrmann (2003) are followed for the purpose of the identification key. In Iberia, *A. nitidula* is considered to be present throughout, but *A. distinguenda* is limited to the extreme north-east of Spain. However, it is clear that there is a major disagreement between the morphological and genetic methods, and these two species may ultimately not be consistently separable. Further analysis using more powerful genetic techniques is necessary.

Relictual species morphologically related to Andrena (Micrandrena) rugulosa Stöckhert, 1935

Inspection of *Micrandrena* specimens from high altitude in the Sierra Nevada revealed the presence of a species that morphologically resembles *A.* (*Micrandrena*) rugulosa Stöckhert, 1935 due to its head that is only slightly shorter than wide rather than clearly shorter than wide, an unusual character in *Micrandrena*. This finding is remarkable, because although widely distributed in Central and Eastern Europe, *A. rugulosa* has a western limit in the Swiss Alps, and has not been previously recorded from France, Spain, or Portugal (Gusenleitner and Schwarz 2002). Samples collected from the Sierra Nevada at high altitude (2000–2100 m) during 2021 are strongly differentiated from *A. rugulosa*, separated by an average genetic distance of 16.01% (range 15.81–16.41%;

Fig. 18). Instead, they form a clade with bootstrap support of 96 with sequences of *A.* (*Micrandrena*) atlantea Wood, 2021 (High and Middle Atlas in Morocco) and a morphologically similar undescribed *Micrandrena* from high altitude (1800–1900 m) in the Moroccan Middle Atlas.

The Sierra Nevada sequences were separated from *A. atlantea* by an average genetic distance of 6.04% (range 5.71–6.38%) and from the undescribed *Micrandrena* by an average genetic distance of 6.34% (range 6.31–6.38%). These three species would therefore seem to represent an isolated *Micrandrena* lineage that is restricted to the Sierra Nevada and the High and Middle Atlas Mountains in Morocco, with consequent genetic and morphological divergence. This is the same pattern as observed in the subgenus *Euandrena*, suggesting that the Sierra Nevada hosts the remnants of a fauna that was presumably once more widespread across Iberia and North Africa. Additional sampling and genetic analysis is required to determine if this pattern holds true for other bee groups. The new *Micrandrena* species from the Sierra Nevada is described below, and the undescribed *Micrandrena* species from Morocco will be described in an upcoming publication.

Andrena (Micrandrena) niveata Friese, 1887 sensu lato

Andrena niveata was described from Germany and Hungary, without further information (Friese 1887). The exact type locality is unclear, as well as the type depository, with no clearly identifiable syntypes located during searches in the SMFD and ZMHB collections, the suggested possible depositories (Gusenleitner and Schwarz 2002). Nevertheless, the concept of A. niveata in Central Europe is clear, and A. niveata is a well-defined species in this region (Schmid-Egger and Scheuchl 1997; Gusenleitner and Schwarz 2002; Amiet et al. 2010). Warncke described three subspecies; A. n. lecana Warncke, 1975 (Spain; locus typicus: Rivas-Vaciamadrid), A. n. haloga Warncke, 1980 (Italy; locus typicus: Lecce), and A. n. bubulca Warncke, 1975 (Turkey; locus typicus: Erzurum); the status of the Italian and Turkish subspecies is not addressed here. Unfortunately, all nine freshly collected A. n. lecana specimens from central and southern Spain (including from the locus typicus) sent for genetic analysis failed to produce sequences. More specific primers will be required to address this question using genetic evidence.

Warncke (1975a) described A. n. lecana extremely briefly, noting that in both sexes the first tergum was finely but clearly shagreened, and finely but more strongly punctured than in A. niveata s. str., with the following terga also more clearly and strongly punctured. He then gave a distribution of A. n. lecana across much of central Spain south to hilly parts of the provinces of Granada (Huéscar) and Almería (Tíjola). In contrast he gives a distribution for A. niveata s. str. of the whole of Iberia, though his distribution maps (Gusenleitner and Schwarz 2002) show a distribution covering eastern, central, and south-eastern Spain. There is therefore the possibility that the two subspecies exist in sympatry, suggesting that they may be distinct.

Examination of material from Spain shows that the two taxa are distinct, but morphological separation is challenging, may not be possible in all cases when old,

abraded, or dirty specimens are available, and is best made with reference to the male genital capsule. Taking male specimens, the shape of the gonostyli are distinctive. In A. niveata s. str., the gonostyli are elongate, with the inner margins of the gonostyli only weakly produced towards the penis valves (Fig. 20A). In contrast, in A. n. lecana, the inner margins of the gonostyli are strongly and clearly produced towards the penis valves (Fig. 20B); the apical spatulate part of the gonostyli are also comparatively shorter, thus the gonostyli appear shorter and more compact in A. n. lecana and longer and more elongate in A. niveata s. str. The structure of the genital capsule of A. niveata s. str. is consistent across Spain to the province of Málaga (the most southerly A. niveata s. str. male examined). Furthermore, the male scutum is shagreened in A. niveata s. str., whereas it is extensively shiny in *A. n. lecana*, and the terga are also more strongly shagreened in A. niveata s. str. compared to A. n. lecana (Fig. 20C, D). Separation of females (recognised by their wide and poorly defined propodeal triangle, Fig. 20E, F, in combination with their depressed tergal margins) is more challenging and may not be possible in all situations. Typically, A. niveata s. str. has the terga shagreened, with punctures on the tergal discs partially disappearing into this shagreenation (Fig. 20G). In contrast, A. n. lecana has the terga shiny, almost without shagreen, with the dense punctures on the tergal discs strong and clearly visible, not disappearing due to the absence of background shagreenation (Fig. 20H). The scutum and scutellum are also more extensively shiny, whereas in A. niveata s. str. these areas usually show at least some shagreenation and are never polished and smooth. The problem comes that some individuals nominally assigned to A. niveata s. str. have partially shing terga, and without genetic sequences it is not possible to say with complete confidence if they belong to A. niveata s. str. or A. n. lecana. Moreover, all Iberian A. niveata s. str. have tergal shagreenation that is weaker than that of A. niveata s. str. populations in Central Europe.

The position is taken here that the strength of tergal shagreenation in *A. niveata* s. str. females is variable across Europe, but that *A. lecana* stat. nov. is a valid species based on the overlapping range in combination with the consistent difference in the shape of the male genital capsule. It has a distribution across steppic parts of central Spain, extending into mountainous areas in south-eastern Spain. Females displaying any level of shagreenation on the tergal discs are considered to represent Iberian populations of *A. niveata* s. str., and females with completely polished tergal discs without a trace of shagreenation represent *A. lecana* (see identification key). Future genetic investigation using more targeted primers will be necessary to confirm this position. Finally, specimens of *A. lecana* from high altitude in the Sierra Nevada show slightly different antennal ratios in the male sex, though the male genital capsule is otherwise identical; this requires further investigation.

Material examined. *Andrena lecana*: Spain: Ribas [Rivas-Vaciamadrid, 40.3503°N, -3.5390°E], 6.v.1908, 1♀, leg. Dusmet, OÖLM (holotype); Carboneras de Guadazaón (Cuenca), 1030 m, 16.v.2009, 1♀, leg. F.J. Ortiz-Sánchez, FJOS; Guadalajara, Alcolea del Pinar, 12.v.2021, 1♀, leg. T.J. Wood, TJWC; Guadalajara, Lupiana, 12.v.2021, 1♀, leg. T.J. Wood, TJWC; Huéscar (Granada), 1900, 1♂, leg. Escalera, OÖLM (paratype); La Cabrilla, Sierra Cazorla (Jaén), 1600 m, 1–3.vi.2022,

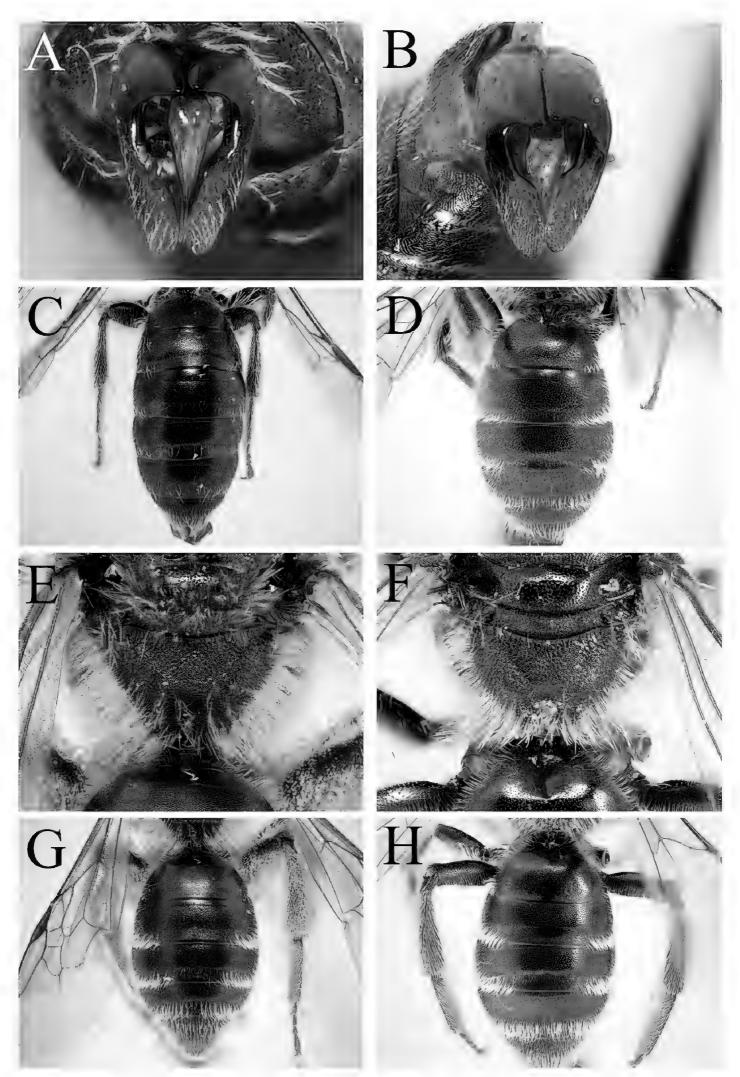


Figure 20. Andrena (Micrandrena) niveata Friese, 1887 **A** male genital capsule **C** male terga, dorsal view **E** female propodeal triangle **G** female terga, dorsal view; Andrena (Micrandrena) lecana Warncke, 1975 **B** male genital capsule **D** male terga, dorsal view **F** female propodeal triangle **H** female terga, dorsal view.

Andrena niveata s. str.: Spain: Cáceres, Cuacos de Yuste, 500 m, 11.v.1999, 1♀, leg. H. & J.E. Wiering, RMNH; Cádiz, Grazalema, Río Guadalete, 28.v.2021, 1♀, leg. T.J. Wood, TJWC; Cádiz, Embalse de Barbate Sw, 1♂, 3♀, 6.v.2017, leg. Barták & Kubik, OÖLM/TJWC; Granada, Sierra de Baza, Prados del Rey, 2000 m, 19.vi.2021, 11♀, leg. T.J. Wood, TJWC; Málaga, 5 km W of Alhaurín de la Torre, 4.vi.1962, 1♀, leg. Jeekel & Wiering, RMNH; Málaga, Cortes de la Frontera, path to Llano de las Labores, 26.v.2021, 3♀, leg. T.J. Wood, TJWC; Málaga, Estepona, 21.iv.1983, 1♀, leg. H. Teunissen, RMNH; Navarra, Tudela, 16.iv.1978, 1♀, leg. C. Gielis, RMNH; Gerona, Figueras, 15.iv.1971, 1♀, leg. J. Leclercq, UMONS.

Andrena (Micrandrena) spreta Pérez, 1895, Andrena (Micrandrena) curtula Pérez, 1903, Andrena (Micrandrena) pusilla Pérez, 1903, and Andrena (Micrandrena) pauxilla Stöckhert, 1935

Members of this species group are challenging to identify and have been inconsistently treated in the literature, with variable species concepts. *Andrena spreta* was described from Algeria, with Warncke (1967) designating a lectotype from Biskra (Fig. 21A; Pérez also mentions a female from Constantine in his catalogue). *Andrena curtula* was described from north-eastern Spain, with Warncke (1967) designating a lectotype from Barcelona (Fig. 21B). *Andrena pusilla* was described from the south of France (Bordeaux, Nantes) and Spain, with Warncke (1967) designating a lectotype from Nantes (Fig. 21C). Finally, *A. pauxilla* was described from the Upper Rhine valley in south-western Germany, with Stöckhert (1935) designating a female from Karlsruhe as the holotype.

Warncke (1967) recognised three species, A. spreta, A. curtula, and A. pauxilla, treating A. pusilla as a subspecies of A. spreta. Warncke (1974) recognised only one taxon in North Africa (A. spreta spreta), but Warncke (1976) surprisingly recognised only A. spreta pauxilla in Iberia, not mentioning A. curtula as a species despite its description from Iberia, only referring to its listing in Ceballos (1956) but indicating that this referred to A. (Micrandrena) tenuistriata Pérez, 1895. This treatment is difficult to understand. Warncke later adopted an extremely broad position (see distribution maps in Gusenleitner and Schwarz 2002), lumping all members of the group into a broad A. spreta. In this model, A. spreta s. str. was restricted to North Africa, A. spreta curtula was distributed across Iberia to southern France, A. spreta pusilla was found in northern and Central Europe, with two more subspecies in i) Italy and ii) south-eastern Europe to the Levant.

This position was not adopted by subsequent authors. Schwarz et al. (1996) accepted *A. pusilla* as a valid species, as well as *A. curtula*. However, they synonymised *A. pauxilla* with *A. curtula*. Schmid-Egger and Scheuchl (1997) did not follow this inter-

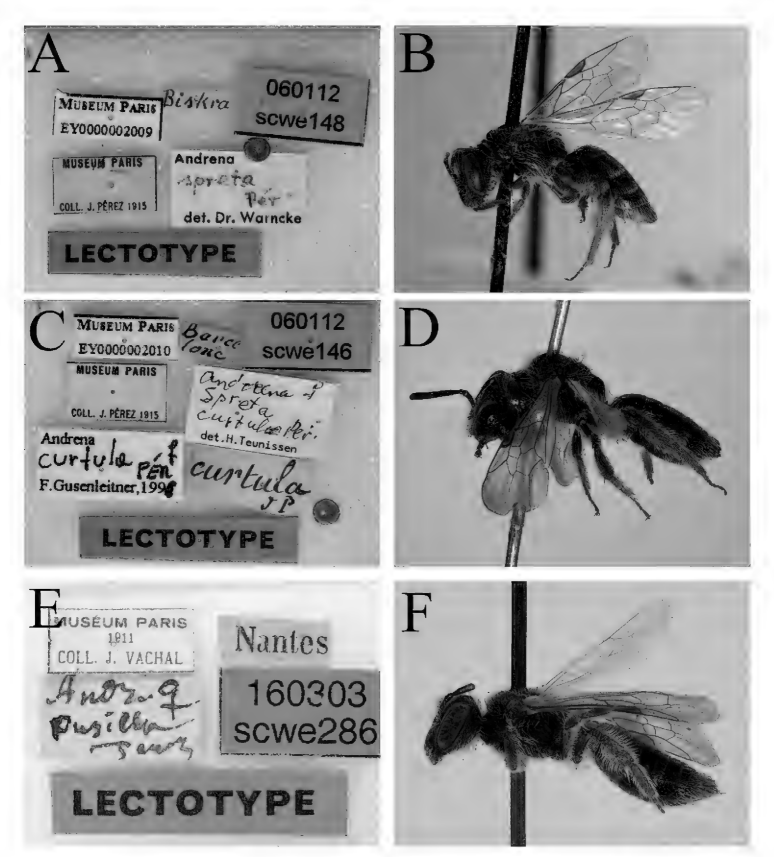


Figure 21. Andrena (Micrandrena) spreta Pérez, 1895, female lectotype A label details B profile; Andrena (Micrandrena) curtula Pérez, 1903, female lectotype C label details D profile; Andrena (Micrandrena) pusilla Pérez, 1903, female lectotype E label details F profile.

pretation, treating *A. pauxilla* as a distinct species, though Amiet et al. (2010) followed the position of Schwarz et al. (1996). Dardón (2010) and Dardón et al. (2014) accepted only a broad *A. spreta* taxon, including *A. curtula*, *A. pusilla*, and *A. pauxilla*, recognising *A. spreta* s. str. and *A. spreta pusilla* in an Iberian context. The situation is therefore unclear, both for the number of species present in this complex, and their distributions.

Analysis of barcodes provides unambiguous support for the existence of three distinct species (Fig. 18), A. spreta including A. curtula, A. pusilla, and A. pauxilla sp.

resurr. Specimens of A. spreta from Israel, Morocco, Portugal, Spain formed a clear clade with bootstrap support of 85. These sequences were separated by an average genetic distance of 0.45% (range 0.00-1.07%). They were clearly separated from a sister clade of A. pusilla sequences from Belgium, France, and Germany by an average genetic distance of 4.33% (range 3.88–4.84%). The A. pusilla clade had low intraspecific genetic distance of 0.29% (range 0.00–0.71%). Andrena pauxilla was strongly differentiated and was found as sister to A. (Micrandrena) rugulosa, showing average genetic differentiation of 10.96% (range 10.38–11.85%) from A. spreta and 11.69% (range 10.75–12.59%) from A. pusilla. This A. pauxilla clade includes a specimen from the south of France (KJ837178, Lac St. Croix) that was identified as A. curtula, the genetic results suggest that it is misidentified. Morphologically, A. pauxilla is clearly differentiated from A. spreta and A. pusilla due to the absence of a gradulus at the base of the terga. Confusion is impossible in the female sex after recognition of this character; the foveae are also longer and narrower and filled with white rather than light brown hairs, the scutum is more densely and uniformly punctate when compared to *A. spreta*. Examination of the lectotypes of A. spreta and A. curtula shows no clear structural difference. The degree of shagreenation varies, but this is typical for A. spreta across its range; both lectotype specimens show the presence of a gradulus at the base of the terga, excluding their conspecificity with A. pauxilla. Due to these genetic results combined with the morphology of the lectotype specimens, the synonymy of A. curtula with A. spreta as proposed by Dardón (2010) and Dardón et al. (2014) is therefore followed, though their synonymies of A. pusilla and A. pauxilla with A. spreta are rejected.

A single barcode was available from Moroccan specimens from the Middle Atlas tentatively identified as *A. pauxilla* which showed an average genetic distance to European *A. pauxilla* specimens of 2.00% (range 1.48%–2.29%). This is considered to represent only separation by distance, and thus *A. pauxilla* is recorded for the first time in North Africa and unambiguously recorded in Spain. Within Spain, *A. pauxilla* appears to be principally recorded from mountain ranges such as the Sierra de las Nieves, the Sierra Nevada (Fig. 22A, B), Sierra Cazorla, Sistema Central, and Sistema Ibérico (Fig. 22C, D). *Andrena pauxilla* would therefore have a currently known distribution of Morocco, Spain, France, and Germany. On the basis of these results, *A. spreta* would appear to be a pan-Mediterranean species, and *A. pusilla* its predominantly northern counterpart (though the species descends south into Italy). The two species can be found in sympatry in the south of France (Praz, in litt.), but to date I have examined no *A. pusilla* specimens from Iberia. Due to the confusion in this group, it must be considered absent until it can be positively demonstrated to be present south of the Pyrenees.

Material examined. *Andrena curtula*: Spain: Barcelona [41.4028°N, 2.1332°E], 1, MNHN (lectotype; Fig. 21B).

Andrena pauxilla: France: Bischenberg, 28.vi.1936, 1♀, leg. M. Klein, det. E. Stöckhert, OÖLM; Hausbergen, 29.vi.1930, 1♀, leg. M. Klein, det. H.R. Schwenninger, OÖLM; Morocco: Fès-Meknès, Boulemane, R503, 7 km SE of Boulemane, 1900 m, 22.v.2022, 1♂, 1♀, leg. T.J. Wood, TJWC; Spain: Canet de Mar, 26.iii.1963, 1♀, leg. F. Vergés, det. H.R. Schwenninger, OÖLM; Cuenca, Huerta del

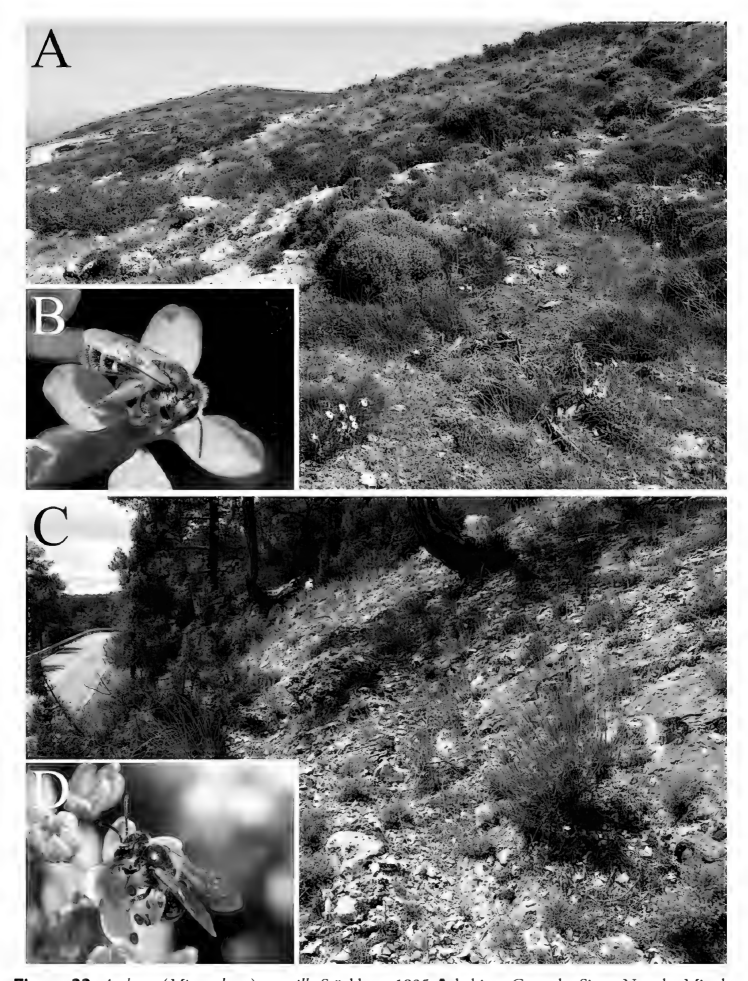


Figure 22. Andrena (Micrandrena) pauxilla Stöckhert, 1935 **A** habitat, Granada, Sierra Nevada, Mirador Monte Ahí de Cara, 2100 m, 12.vi.2021 **B** female collecting pollen from *Vella spinosa* (Brassicaceae) **C** habitat, Cuenca, Mirador Valle de Valdecabras, 21.vi.2021 **D** female collecting pollen from *Sedum* spp. (Crassulaceae).

Marquesado, environs north of town, 26.vi.2021, 3♀, leg. T.J. Wood, TJWC; Granada, Sierra Nevada, Jardín Botánico Hoya de Pedraza environs, 1900 m, 9.vi.2021, 1♀, leg. T.J. Wood, TJWC; Granada, Sierra Nevada, Mirador Monte Ahí de Cara,

2100 m, 12.vi.2021, 1♀, leg. T.J. Wood, TJWC; Guadalajara, Aldeanueva de Atienza, 9.vii.2021, 1♀, leg. T.J. Wood, TJWC; Málaga, PN Sierra de las Nieves, mountain peak S of Pinsapo Escalereta, 30.v.2021, 1♀, leg. T.J. Wood, TJWC; La Cabrilla, Sierra Cazorla (Jaén), 1600 m, 3.vi.2022, leg. C.M. Herrera, CMHC; Sierra Cazorla, Puerto Llano, 1800 m, 11.vi.2022, 1♂, 3♀, leg. J. Valverde, CMHC; Teruel, Guadalaviar, Rambla de los Ojos, 27.vi.2021, 1♀, leg, T.J. Wood, TJWC; Teruel, Villar del Cobo, Barranco de los Oncenachos, 27.vi.2021, 6♀, leg, T.J. Wood, TJWC; Cuenca, Mirador Valle de Valdecabras, 21.vi.2021, 2♀, leg, T.J. Wood, TJWC.

Andrena pusilla: France: Nantes [47.2233°N, -1.5542°W], 1, MNHN (lectotype; Fig. 21C).

Andrena spreta: ALGERIA: Biskra [34.8600°N, 5.6995°E], 1\,\tilde{2}, MNHN (lectotype; Fig. 21A).

Andrena (Micrandrena) strohmella Stöckhert, 1928 and Andrena (Micrandrena) icterina Warncke, 1974

Andrena strohmella was described from southern Germany and is a typically early spring species in the Central European Andrena fauna, with records extending south to the High and Maritime Alps in France, and west to the Bordeaux region; it has not previously been reported from the Pyrenees (Gusenleitner and Schwarz 2002). Warncke (1974) later described A. icterina from northern Algeria. He later indicated the presence of this species in south-eastern Spain in his distribution maps (Gusenleitner and Schwarz 2002), though he did not list its present in Iberia (Warncke 1976). Wood et al. (2020b) later reported the presence of this species in northern Morocco. In his description of A. icterina, Warncke draws comparison with A. strohmella, noting that the process of the labrum is half as broad, the clypeus is more sparsely punctate and lacking an unpunctured midline, that the terga are more strongly shagreened, and that the carinae on the dorsolateral corners of the first tergum are more weakly pronounced. Dardón (2010) and Dardón et al. (2014) did not actually examine any material of Iberian A. icterina, but inspection of new Iberian material has demonstrated that A. icterina is more widespread in Iberia than previously thought, being found from the Los Alcornocales to the Sierra de las Nieves, the Sierra Nevada, Sierra de Baza, Sierra Cazorla, and Sistema Central, and is also newly reported from northern Portugal (see below). In all of these localities it is found in mountainous areas, generally at high altitude in excess of 1000 m.

Dardón (2010) and Dardón et al. (2014) reported *A. strohmella* from north-east-ern Spain, from Tarragona: Alcanar, v.2002, leg. Kadlec, OÖLM. I have not been able to locate and examine this specimen for myself in the OÖLM collection, but it is slightly problematic for ecological regions. *Andrena strohmella* is clearly a species of deciduous temperate woodland; this is reflected in its distribution across Central Europe, and in its association with flowering trees such as *Crataegus* (Rosaceae), *Prunus* (Rosaceae), and *Salix* (Salicaceae). *Andrena strohmella* has therefore not previously been reported from Mediterranean regions. The record from Alcanar does not therefore match the ecology of this species, Alcanar being a low-elevation (c. 70 m) village close

to the Mediterranean coast. However, *A. strohmella* is present in north-eastern Catalonia based on recently collected specimens (Álvarez Fidalgo, in litt.), and so the species is retained on the Spanish and Iberian lists.

More broadly, morphological differences between *A. strohmella* and *A. icterina* are slight, though they do not appear to introgress based on examined specimens. Some characters such as the strength of the carinae on the dorsolateral corners of the first tergum are not completely consistent, because some specimens in Central Europe can be found in which these are very weakly produced. Genetic data will hopefully clarify the status of *A. icterina*, but unfortunately, like *A. lecana*, this taxon appears to be challenging to barcode, as all seven Iberian specimens sent for genetic analysis failed or returned corrupted sequences.

Material examined. Andrena icterina: Portugal: Bragança, Serapicos, 16.v.2021, 12, leg. A. Soares, A. Soares Coll.; **Spain:** Ávila, Hoyocasero, El Pinar de Hoyocasero, 16.v.2021, 1♀, leg. T.J. Wood, TJWC; Ávila, Navalsauz, 1 km E, Alberche stream, 16.v.2021, 1♀, leg. T.J. Wood, TJWC; Cádiz prov., Vent. L. Canillas Hozgarganta-Tal b. Jimena 250 m, 14.iv.1985, 32, leg. W. Schacht, OÖLM; Campamento Alfaguara (Alfacar, Granada), 1420 m, 13.v.2007, 13, 22, leg. F.J. Ortiz-Sánchez, FJOS; Cortijo Tortas, Paterna del Madera (Albacete), 1310 m, 30.iv.2022, 26, leg. F.J. Ortiz-Sánchez, FJOS; Granada, Sierra de Baza, Prados del Rey, 2000 m, 19.vi.2021, 12, leg. T.J. Wood, TJWC; Granada, Sierra Nevada, Capileira to La Cebadilla, 1500 m, 8.vi.2021, 12, leg. T.J. Wood, TJWC; Granada, Sierra Nevada, Jardín Botánico Hoya de Pedraza environs, 1900 m, 2\, leg. T.J. Wood, TJWC; Granada, Venta de los Alazores, 25.v.1982, 1, leg. R. Leys, RMNH; La Cabrilla, Sierra Cazorla (Jaén), 1600 m, 3.vi.2022, leg. C.M. Hererra, CMHC; Málaga, PN Sierra de las Nieves, mountain peak S of Pinsapo Escalereta, 1600 m, 30.v.2021, 32, leg. T.J. Wood, TJWC; Sierra Cazorla, Puerto Llano, 1800 m, 11.vi.2022, 12, leg. J. Valverde, CMHC; Puerto Crucetillas, Riópar (Albacete), 1340 m, 30.iv.2022, 13, leg. F.J. Ortiz-Sánchez, FJOS; Sierra de Cazorla, Nava de las Correhuelas, 1.v.2021, 13, leg. C.M. Hererra, CMHC.

Subgenus Notandrena Pérez, 1890

The definition of this subgenus was expanded by Pisanty et al. (2022b) to include *Carandrena* Warncke, 1968 (type species *Andrena aerinifrons* Dours, 1873). It is important to note that several species formerly placed within the *Carandrena* fall elsewhere into an undescribed subgenus, but these are desert-living species, none of which are present in Iberia. In the context of this revision, *Carandrena* is treated as a strict synonym of *Notandrena*.

Andrena (incertae sedis) urdula Warncke, 1965 was described from Greece and is a rare and poorly understood taxon known only from a small number of specimens. It is reliably known only from Greece (type series), Spain (central Spain), and Morocco, as the distribution map presented by Gusenleitner and Schwarz (2002: 1201) actually refers to A. (Notandrena) ungeri Mavromoustakis 1952; the distribution map for A. ungeri (p. 1200) is actually the distribution map for A. urdula. A barcoded specimen

from south of Madrid [WPATW351-21] does not clearly fall into the *Notandrena* group (Fig. 23); its true placement is unclear because it does not have any strong characters that lead to obvious affinities with specific groups. It is best treated as incertae sedis until more genetic data are available.

Warncke described two similar taxa in the group of small metallic green *Notandrena*, *A.* (*Notandrena*) reperta Warncke, 1974 and *A.* (*Notandrena*) reperta varuga Warncke, 1975. Warncke actually described *A. reperta* as a subspecies of *A. varuga*, but due to the order of publication, *A. reperta* has priority. Both species can be recognised because the hind tibial spur is apically curved, though this character is more pronounced in Iberian specimens. Genetically, specimens of *A. reperta* from Morocco and *A. reperta varuga* from central Spain formed a clade with a specimen of *A.* (*Notandrena*) nigroviridula Dours, 1873 from Morocco. All three taxa were well separated; *A. reperta* and *A. reperta varuga* by 12.84%, *A. reperta* and *A. nigroviridula* by 11.67%, and

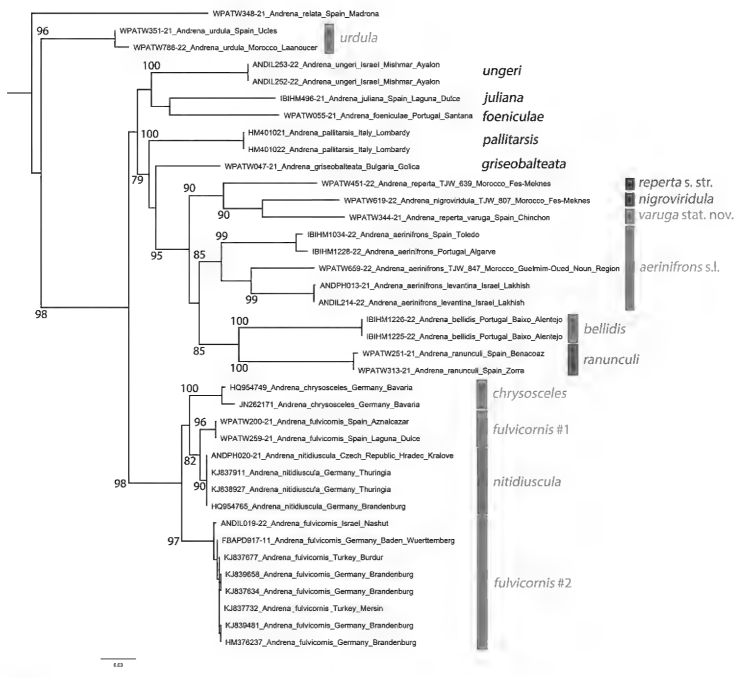


Figure 23. Phylogenetic tree (maximum likelihood) of *Andrena* subgenus *Notandrena* Pérez, 1890 based on the mitochondrial COI gene. *Andrena* (incertae sedis) *relata* Warncke, 1975 is used as an outgroup. Numbers adjacent to branches represent bootstrap support (values of <75 are omitted).

A. reperta varuga and A. nigroviridula by 10.12% (Fig. 23). Moreover, A. reperta and A. reperta varuga do not form a monophyletic clade, being rendered paraphyletic by A. nigroviridula. On this basis, A. varuga stat. nov. is raised to species status. Andrena varuga is endemic to Spain, and A. reperta is found in Morocco, Algeria, and Tunisia.

Dours (1873) described A. aerinifrons from Algeria, and A. aerinifrons levantina Hedicke, 1938 was later described from the Levant. Barcode analysis shows that specimens from Israel, Morocco, Portugal, and Spain show high average intraspecific variation of 8.64% (range 0.00–11.67%). The Moroccan specimen is intermediate between the Iberian and Levantine sequences, being separated from the Iberian sequences by an average genetic distance of 9.92% (range 9.73–10.12%) and from the Levantine sequences by 10.12% (range 10.12–10.12%). Given that these sequences form a monophyletic clade, a broad interpretation of A. aerinifrons is taken here; a neotype can be designated for A. aerinifrons at a later date when the situation becomes clearer, as Dours' collection was destroyed in a fire and is not available for study (see below).

Andrena (Notandrena) fulvicornis Schenck, 1861 has been recognised as distinct from A. (Notandrena) nitidiuscula Schenck, 1853 (Schmid-Egger and Doczkal 1995; Schmid-Egger and Scheuchl 1997; Schwenninger 2013; Bénon and Praz 2016). Barcode analysis shows that specimens from Germany, Israel, and Turkey are indeed clearly distinct from A. nitidiuscula, not forming a sister clade and showing average separation of 3.79% (range 3.50–3.89%), with bootstrap support of 96. However, sequences of putative A. fulvicornis from Spain fall into a clade of A. nitidiuscula and A. (Notandrena) chrysosceles (Kirby, 1802). Spanish specimens are separated from A. nitidiuscula by 1.95% (range 1.95–1.95%), from A. chrysosceles by 4.67% (range 4.28–5.06%), and from the other clade of *A. fulvicornis* by 4.18% (range 3.89–4.28%). This Iberian 'fulvicornis' is also well-supported, with bootstrap support of 96. At the present time, no taxonomic action can be taken until more genetic data are available. Morphologically, Iberian 'fulvicornis' do not appear to be different from Central European/Turkish/Levantine specimens. Additional sequences from Iberia and France are needed; this barcode difference may be the result of an isolated population in the Iberian glacial refugium.

Subgenus Plastandrena Hedicke, 1933

Pisanty et al. (2022b) synonymised the subgenus *Agandrena* with *Plastandrena*, and this broader definition is used here. Some taxonomic problems within this subgenus were discussed by Wood (2023a), who found support for the species-status of *A. (Plastandrena) nigrospina* Thomson, 1872. Standard barcodes have limited utility for this group for unclear reasons, so care must be taken when interpreting results, as morphologically distinct taxa can form unclear clades, for example *A. (Plastandrena) tibialis* (Kirby, 1802). This result was reproduced here (Fig. 9), and no further discussion or action is taken; in an Iberian context, members of this subgenus appear to be identifiable using barcodes, but work is needed to revise this group at a Palaearctic scale using more powerful genetic techniques.

Andrena (Plastandrena) pilipes Fabricius, 1781 and Andrena (Plastandrena) nigrospina Thomson, 1872

Ortiz-Sánchez et al. (2022) reported A. nigrospina as new for Spain from the Sierra de Cazorla based on male specimens with their distinctive genital capsule. Wood (2023a) further reported two barcoded specimens from Spain (Sistema Central and Sistema Ibérico) belonging to A. nigrospina. This genetic result was further supported with the addition of more sequences from Morocco (Fig. 9), revealing that A. nigrospina is also present in the Middle Atlas. The A. nigrospina clade maintained a low intraspecific genetic distance of 0.41% (range 0.00-1.06%) from Morocco to Kyrgyzstan, and average separation from A. pilipes was 1.63% (range 1.06–2.35%). As this difference is still small in absolute terms, bootstrap support was moderate, with 77 for A. pilipes and 84 for A. nigrospina, but both are considered to be distinct species. Females cannot currently be consistently separated morphologically, so no characters are given in the identification key. In Iberia, A. nigrospina is newly reported for northern Portugal from close to the Peneda-Gerês National Park. The species is likely to be restricted to cooler parts of northern, central, and eastern Iberia where it will probably be found only in mountainous regions, or at least at elevation. More sampling and genetic study is required to define its exact range limits.

Material examined. *Andrena nigrospina*: Portugal: Minho, Ruivães, N103, 12.v.2019, 1♂, 1♀, leg. Wood, TJWC; Spain: Cuenca, Pajaroncillo, 3 km SW, Arroyo de Peña Quebrada, 26.vi.2021, 4♀, leg. T.J. Wood, TJWC (barcoded); Guadalajara, Bustares, 2 km N, Alto Rey, 1780 m, 1♀, 9.vii.2021, leg. T.J. Wood, TJWC (barcoded).

Andrena (Plastandrena) agilissima Scopoli, 1770 and Andrena (Plastandrena) asperrima Pérez, 1895

Andrena agilissima is a widespread West Palaearctic species that is well-known in Central and Southern Europe to north-western Africa. In contrast, A. asperrima is much less well known, having a more Mediterranean distribution in France, Spain, Morocco, Algeria, and Tunisia. Unlike A. agilissima, A. asperrima is bivoltine and is exceptionally variable in the density, size, and strength of the integumental punctation. In the female sex, the typical form has strong and dense punctures on the terga, allowing easy separation from A. agilissima in which the terga have small and subtle punctures. However, many specimens of A. asperrima can be found which have greatly reduced tergal punctation and which are therefore extremely similar to A. agilissima; they can be separated by the smaller body size and the sparser punctation of the scutum. This sparsely punctate form is more common in the south-west and was described from Morocco by Warncke as A. asperrima alascana Warncke, 1974. Overall, the two species are clearly separable by their genital capsules in the male sex.

Because of this variation, it is important to ensure that Iberian material is conspecific with North African material, since the oldest names of Pérez (1895) were described based on North African material. Warncke (1967) designated a male lectotype for

A. asperrima using a specimen from Biskra from the collection of Pic (Fig. 24A, B). This is unjustified, as it is not part of the original syntypic series, and Warncke's designation is here rejected. In Pérez's catalogue, under entry '1030 Andrena trachodes J.P.' [an unpublished name; Pérez sometimes changed his mind and therefore a different name to the published name can be present in the catalogue] Pérez writes that the species comes from Constantine [in Algeria], drawing comparison with A. flessae Panzer, 1805 (= A. agilissima (Scopoli, 1770)) and arguing it differs by the stronger punctation of the metasoma. Inspection of the Pérez collection shows the presence of a female

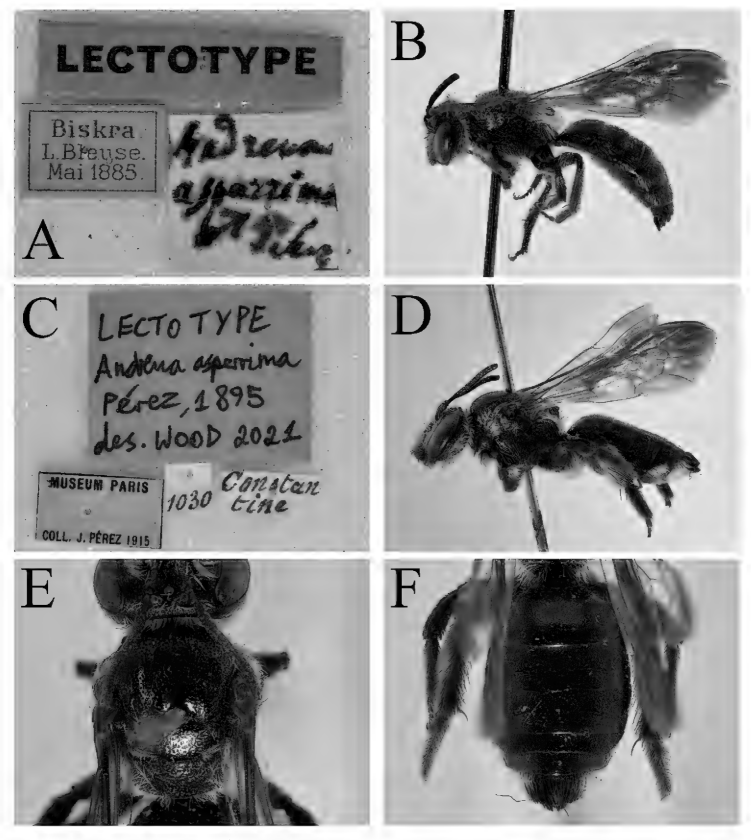


Figure 24. Andrena (Plastandrena) asperrima Pérez, 1895, false male lectotype **A** label details **B** profile; true female lectotype **C** label details **D** profile **E** scutum, dorsal view **F** terga, dorsal view.

specimen from Constantine which bears the label '1030', the code used by Pérez for this species. This specimen is designated as a new lectotype (Fig. 24C–F); it conforms to the concept of *A. asperrima* used by subsequent authors.

Andrena atricapilla Pérez, 1895 was also described from Algeria, but only in the male sex. Warncke (1967) listed this taxon as a synonym of *A. asperrima*, though he did not mention a lectotype. A male specimen labelled as atricapilla J.P. in the handwriting of Pérez is separated in the MNHN collection and labelled as a lectotype, probably by Teunissen as one of his determination labels is present. This lectotype designation was never published, and so it is here designated as a lectotype (Fig. 25). The synonymy with *A. asperrima* is maintained, as the genital capsule is typical for the species concept.

Genetic analysis of specimens of *A. agilissima* and *A. asperrima* from Croatia, Morocco, Portugal, and Spain showed two clear clades (Fig. 9). One clade contained only specimens determined as *A. agilissima* from Croatia, Portugal, and Spain. These sequences showed average intraspecific variation of 0.07% (range 0.00–0.25%), and were strongly separated from the second clade by an average genetic distance of 9.33% (range 8.74–10.20%). This second clade contained specimens identified as *A. agilissima* from the Middle Atlas. There can be no doubt as to the identity of these specimens, as two are males, and their genital capsule is that of *A. agilissima* and not *A. asperrima*. However, the genetic differentiation within this clade is very low at an average of 0.42% (range 0.00–0.91%). As for *A. bimaculata* and *A. tibialis*, there is clearly genetic complexity here, as two species with distinct genital capsules have barcodes that form intermixed clades. No taxonomic action is taken on the basis of these barcoding results.

However, what can be concluded is that Iberian material of *A. asperrima* is conspecific with North African material, including the weakly punctate form that dominates in Morocco. In combination with the new lectotype designations, the invariant genital capsule, and these genetic results, the concept of Warncke (1967; 1974) and Gusenleitner and Schwarz (2002) regarding *A. asperrima* is maintained. The full synonymy is given below.

Andrena (Plastandrena) asperrima Pérez, 1895

Andrena (Plastandrena) asperrima Pérez, 1895: 33, ♀♂ [Algeria, lectotype by present designation: MNHN].

Andrena (Plastandrena) atricapilla Pérez, 1895: 33, & [Algeria, lectotype by present designation: MNHN].

Andrena (Plastandrena) flessae var. elcheensis Friese, 1922: 211, ♀ [Spain: ZMHB, not examined].

Andrena (Plastandrena) hemicyanea Cockerell, 1930: 112, ♀ [Tunisia: type lost?] Andrena (Plastandrena) asperrima alascana Warncke, 1974: 36, ♀♂ [Morocco: OÖLM, examined].

Distribution. Spain, France, Morocco, Algeria, Tunisia.



Figure 25. Andrena (Plastandrena) atricapilla Pérez, 1895, male lectotype **A** label details **B** profile **C** terga, dorsal view **D** genital capsule.

Material examined. ALGERIA: Constantine [36.3645°N, 6.6409°E], 1♀, MNHN (lectotype of *A. asperrima*, by present designation; Fig. 24C–F); Biskra, v.1885, 1♂, leg. Blause, MNHN (false lectotype of *A. asperrima*; Fig. 24A, B); Biskra [34.8600°N, 5.6995°E], 1♂, MNHN (lectotype of *A. atricapilla*, by present designation; Fig. 25); MOROCCO: Tizi-n-Talrhemt nr. Midelt [32.6821°N, -2.9344°E], 1900 m, 1.vi.1968, leg. M.A. Lieftinck, OÖLM (holotype of *A. asperrima alascana*).

Subgenus Poecilandrena Hedicke, 1933

This subgenus is strongly polyphyletic (Pisanty et al. (2022b), but these classification issues relate to eastern taxa; in Iberia, the species belong to *Poecilandrena* s. str. Warncke (1976) listed only one *Poecilandrena* species in Iberia, *A.* (*Poecilandrena*) labiata Fabricius, 1781. Ceballos (1956) and Pérez-Íñigo (1984) also listed *A.* (*Poecilandrena*) potentillae Panzer, 1809, though this species was not listed as present by Warncke and his distribution maps (Gusenleitner and Schwarz 2002) indicate that this species has a western range limit in central France.

Collection of material from the Sierra Nevada produced red-marked *Poecilandrena* females [WPATW281-21] that morphologically resemble *A. potentillae* in the reduced

punctation density at the edge of the clypeus. No 'potentillae' males with their distinctive genital capsule could be found. A female sequence clearly fell into a clade with an A. labiata sequence from Belgium, the two specimens separated by 2.87% (Fig. 1). The four A. labiata sequences from Belgium, Portugal, and Spain showed average intraspecific variation of 2.52% (range 0.78–3.92%), much lower than the average separation from A. potentillae sequences from Belgium, France, Germany, and Hungary of 11.81% (range 11.23–12.27%). The position is taken here that true A. potentillae is not present in Iberia, even though some females of A. labiata can resemble it morphologically. These specimens conform to the subspecific concept of A. labiata bellina Warncke, 1967 that was described from Madrid. Warncke (1967) noted that this form had weaker shagreenation and finer punctation than the nominate form, and I believe that this is the source of the confusion and the erroneous reports of A. potentillae in Iberia. Andrena potentillae is therefore not included in the identification key as typical female characters that can be used to separate the two species in Central Europe do not appear to work universally in Iberia.

Finally, Ortiz-Sánchez (2011, 2020) lists A. (Poecilandrena) viridescens Viereck, 1916 as part of the Spanish fauna. The distribution maps of Warncke (Gusenleitner and Schwarz 2002) indicate the possible presence of this species on the Spanish side of the Pyrenees. I have not examined any specimens of A. viridescens from Iberia, but its presence in the Pyrenees is plausible and so it is retained on the list. Several Andrena taxa with apparent range limits in the Pyrenees have recently been confirmed to occur in northern Spain (e.g. A. (Ulandrena) polita Smith, 1847 and A. (Micrandrena) nanula Nylander, 1848, see Álvarez Fidalgo et al. 2022), and so additional searches may well confirm the presence of A. viridescens as well.

Subgenus Simandrena Pérez, 1890

Barcode analysis returned *Simandrena* as paraphyletic (Fig. 4), but this means very little, as the subgenus is very well characterised morphologically and genetically based on UCE analysis (Pisanty et al. 2022b). No major changes in *Simandrena* taxonomy are made here, but there are a number of issues to discuss.

Warncke (1967) described A. (Simandrena) combinata crudelis Warncke, 1967 from Spain. Comparison of sequences shows that Iberian material is only weakly differentiated, being separated from A. (Simandrena) combinata (Christ, 1791) sequences from Germany by an average genetic distance of 1.12% (range 1.06–1.29%). Based on this evidence, a subspecific status is not justified.

Andrena (Simandrena) vetula Lepeletier, 1841 was recently placed in the Simandrena, as its unusual male morphology had led to confused previous placement (Pisanty et al. 2022b). Genetically there was a large difference between Iberian and North African sequences, separated by an average genetic distance of 6.33% (range 6.08–6.57%). Andrena vetula has an enormous range, from Morocco and Iberia to Central Asia (Wood and Monfared 2022). Sequences are required over this range before any taxonomic changes can be made, as morphologically there are no obvious differences between Iberian and North African specimens.

The species pair of A. (Simandrena) confinis Stöckhert, 1930 and A. (Simandrena) congruens Schmiedeknecht, 1884 continues to pose problems. Warncke (1967) treated A. confinis as a synonym of A. congruens, but others have not followed this interpretation. The two taxa can be separated morphologically in Central Europe (Schmid-Egger and Scheuchl 1997), and there are also ecological differences with A. confinis preferring cooler northern climates and A. congruens preferring warmer and drier southern climates, with overlap in Central Europe. However, in southern Europe the situation is complex and confused. Wood et al. (2021) added A. confinis to the Iberian list on the basis of a specimen from the Picos de Europa Mountains. Genetically, this specimen clusters with A. confinis sequences from Germany. However, sequences from specimens provisionally identified as A. congruens from northern Portugal as well as a specimen of A. congruens from Bulgaria render A. congruens paraphyletic. The genetic distance between all sequences is low. The Portuguese specimens are strongly divergent morphologically from the Spanish A. confinis specimen, with dense and obvious tergal punctation compared to terga which are almost impunctate, but the average genetic differentiation is low at 0.88% (range 0.60–1.15%). No major taxonomic decision is made here; much more genetic data are required for a dedicated study focused on this group. Both nominal taxa are included in the identification key.

Andrena (Simandrena) cilissaeformis Pérez, 1895, sp. resurr.

Andrena (Simandrena) cilissaeformis Pérez, 1895: 42, ♀ [Spain, lectotype by present designation: MNHN]

Andrena breviscopa auctorum.

Remarks. Andrena breviscopa Pérez, 1895 was described in the female and male sexes from North Africa. Warncke's treatment of A. breviscopa is curious, because he designated a lectotype (Fig. 26) and listed the taxon as a synonym of A. numida Lepeletier, 1841 (Warncke 1967). However, just a few years later he listed A. breviscopa as a valid taxon in the subgenus Simandrena (Warncke 1974). Examination of the female lectotype designated by Warncke shows that his original synonymy was correct; the taxon is clearly not a Simandrena, and is indeed a synonym of A. numida.

The use of the name *A. breviscopa* to apply to the taxon present in Spain, Morocco, and Algeria is therefore incorrect. The correct name is *A. cilissaeformis* Pérez, 1895 sp. resurr. *Andrena cilissaeformis* was described from Spain, not Algeria as stated in Warncke (1967) and Gusenleitner and Schwarz (2002). This is because Pérez (1895) does not directly state the type locality (or indeed, any information about the collecting locality of any of the species described in this work), but this information is included in his personal catalogue. Warncke (1967) did not examine material of *A. cilissaeformis*, stating that whilst the label was present, material was missing. Examination of material in the MNHN has located a specimen labelled with 'Esp' [Espagne = Spain] in Pérez's handwriting (Fig. 27). This specimen was separated by Teunissen, but he never published this information. This specimen is badly damaged, but it is a *Simandrena* and

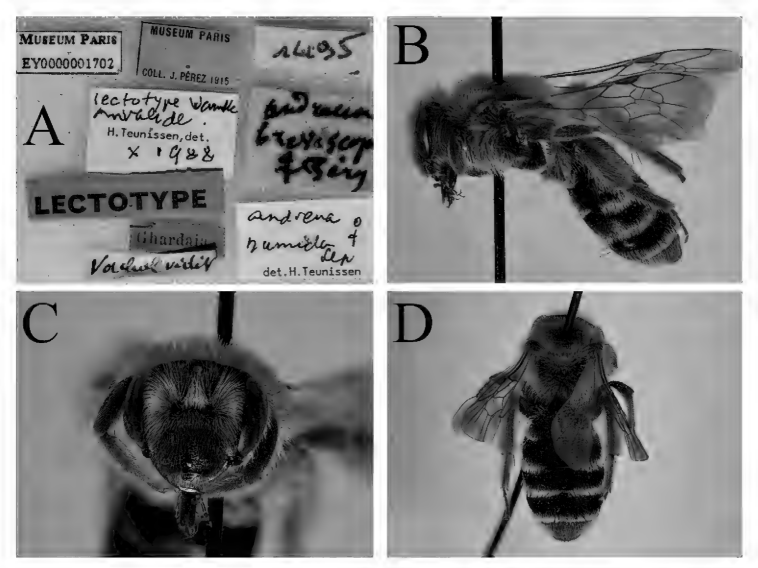


Figure 26. *Andrena* (incertae sedis) *breviscopa* Pérez, 1895, female lectotype **A** label details **B** profile **C** face, frontal view **D** terga, dorsal view.

conforms to *A. breviscopa* auctorum sensu Warncke (1974) and subsequent publications. It is designated as a lectotype to fix the name on the Iberian population. *Andrena breviscopa* is returned to its original synonymy with *A. numida*.

Distribution. Spain, Morocco, Algeria.

Material examined. *Andrena breviscopa*: ALGERIA: Ghardaia [32.5047°N, 3.6419°E], 1, MNHN (lectotype; Fig. 26); (*Andrena cilissaeformis*): **SPAIN:** no collection information, 1, MNHN (lectotype by present designation; Fig. 27).

Subgenus Taeniandrena Hedicke, 1933

Large parts of this subgenus have been revised recently by Wood et al. (2021), Praz et al. (2022), Wood (2022), and Wood and Ortiz-Sánchez (2022). There is relatively little new information to present here, other than to revise the status of *A.* (*Taeniandrena*) *poupillieri* Dours, 1872, and so a reduced phylogenetic tree is presented given the results presented in these previous publications (Fig. 28). However, it is clear that there is still unfinished work to be completed in this subgenus in an Iberian context. Specifically, Praz et al. (2022) identified an unclear lineage "sp. nov. 2" from northern Portugal. This specimen is a male that has a genital capsule that diverges from any known Iberian

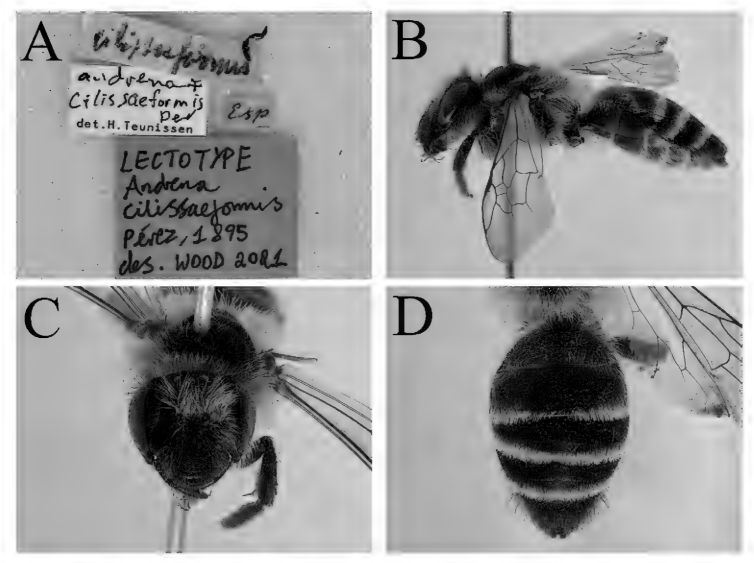


Figure 27. Andrena (Simandrena) cilissaeformis Pérez, 1895; female lectotype A label details B profile C face, frontal view D terga, dorsal view.

species. Furthermore, I have an unbarcoded male from Guadalajara province that also possesses a genital capsule that does not match any known species. These may represent additional undescribed *Taeniandrena* species endemic to the peninsula. No further action is taken until more genetic samples are available; it is extremely likely that additional barcoding will discover more diversity in this challenging and speciose subgenus.

Andrena (Taeniandrena) poupillieri Dours, 1872

Two further issues require discussion. The first is the identity of *A. poupillieri*. Praz et al. (2022) identified two potential mitochondrial lineages that could correspond to this species. Additional sampling in Morocco has clarified the situation; *poupillieri 1* sensu Praz et al. (2022) corresponds to *A. (Taeniandrena) gregaria* Warncke, 1974 and *poupillieri 2* sensu Praz et al. (2022) corresponds to the concept of *A. poupillieri* used by Warncke. As the type of *A. poupillieri* is lost, it is beneficial to designate a neotype in order to fix the concept of this species in line with the existing literature; a specimen from Algeria is chosen, as this is the original *locus typicus*. As *A. gregaria* does not occur in Iberia, it is not included in the phylogenetic tree presented here; it will be dealt with in the upcoming North African *Andrena* revision. The species *A. poupillieri* is

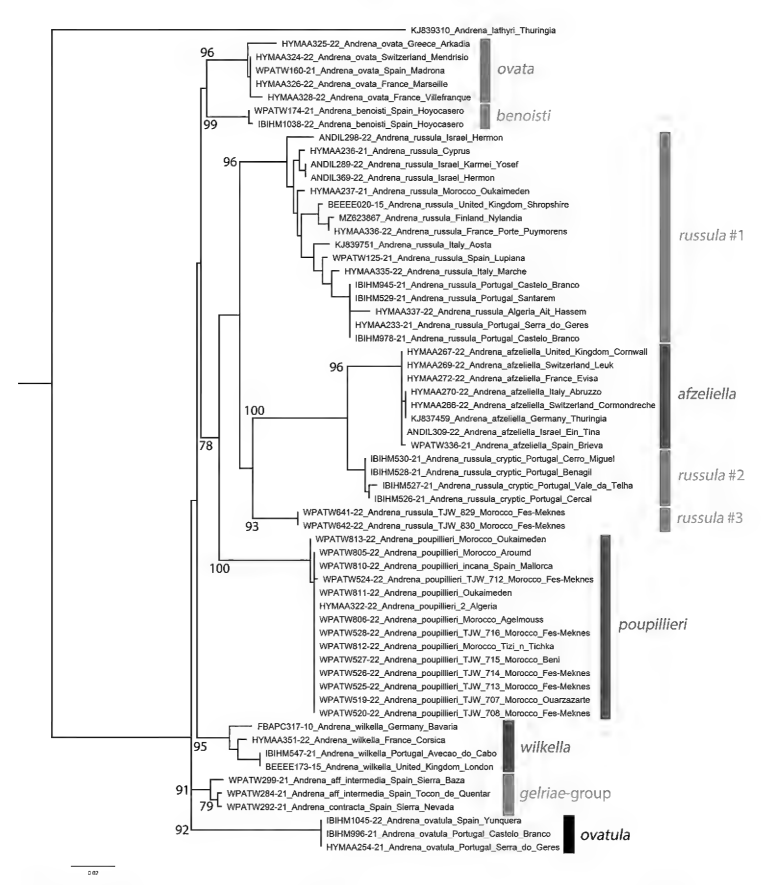


Figure 28. Phylogenetic tree (maximum likelihood) of *Andrena* subgenus *Taeniandrena* Hedicke, 1933 based on the mitochondrial COI gene. *Andrena* (*Taeniandrena*) *lathyri* Alfken, 1900 is used as an outgroup. Numbers adjacent to branches represent bootstrap support (values of <75 are omitted).

commonly encountered in Morocco, and is clearly identifiable from barcodes, forming a clade with bootstrap support of 100. Separation of females from *A. (Taeniandrena) ovatula* (Kirby, 1802) can be made by the tergal punctation, this being much stronger in *A. ovatula*. Generally, the two taxa are well-separated by an average genetic distance of 6.12% (range 5.86–7.79%). Males of *A. poupillieri* can be recognised based on the genital capsule, in which the gonostyli are apically produced into acute points.

Warncke changed his mind about the status of *A. poupillieri* – in Warncke (1967), he lists the taxon as A. ovatula poupillieri, but then in Warncke (1975a) he described A. poupillieri incana from the Balearic islands, listing A. poupillieri s. str. from southern Iberia (Warncke 1976). In his distribution maps (Gusenleitner and Schwarz 2002), he gives a distribution of southern Iberia, the Balearic islands, north-western Africa, Libya, and Crete. Records from Crete are likely to refer to unrecognised A. (Taeniandrena) ovata Schenck, 1853 (Wood, unpublished data). Examination of male specimens from the extreme south of Spain shows that A. poupillieri is present based on the distinctive genital capsule, though as females cannot be separated morphologically from A. ovatula the exact range of the two species is unclear. Andrena ovatula reaches at least as far south at the Sierra de las Nieves in Málaga province [IBIHM1045-22], though this far south it may be restricted to mountainous areas whereas the two examined male A. poupillieri specimens come from the coast. More collection is required. For now, A. poupillieri is considered to be a rare and little-collected taxon in Iberia, probably restricted to the coast in the south and south-east of the peninsula. A single sequence for A. p. incana was available from Mallorca that unambiguously nests within the North African A. poupillieri sequences; it is identical to 12 of the North African sequences, and differs from the thirteenth by just 0.15%. As such, A. poupillieri including A. p. incana shows extremely low intraspecific variation of 0.03% (range 0.00-0.15%), and subspecific status is unnecessary for the population on the Balearic islands.

Material examined. Spain: Málaga, Estepona, 24.iii.1986, 1&, leg. J. van Oosterhout, RMNH; Málaga, San Pedro de Alcántara, 15.iii.1986, 1&, leg. C. v. Achterberg, RMNH.

Andrena (Taeniandrena) russula Lepeletier, 1841

Praz et al. (2022) synonymised A. (Taeniandrena) similis Smith, 1849 with A. russula, taking a broad, pan-Mediterranean approach. In an Iberian context, sequences from central and northern Iberia clearly fall into a broad A. russula clade along with sequences from Morocco to Cyprus and Israel and north to the United Kingdom and Finland (Fig. 28). However, in the extreme south-west of Iberia in southern Portugal, sequences from specimens that are morphologically indistinguishable from A. russula from the rest of Iberia form a distinct clade that falls closest to A. (Taeniandrena) afzeliella (Kirby, 1802). These specimens come from the south of Baixo Alentejo (Cercal) and the Algarve (Aljezur, Benagil, Moncarapacho). These sequences are consistently separated from the broad A. russula clade by 5.50% (range 3.68–11.11%). Two specimens from Morocco that correspond to A. russula form a third clade of A. russula s.l. At the present time, no taxonomic action is taken; these three lineages may represent different isolated populations. More powerful genetic techniques are required to resolve this problem, as for Iberian members of the gelriae-group (see Praz et al. 2022).

Andrena (Taeniandrena) gredana Warncke, 1975

Wood et al. (2021) elevated *A. gredana* to species status, and gave a distribution across the Sistema Central, central and northern Portugal, and northern Spain across to the

Pyrenees in the province of Huesca. Examination of material from the Hautes-Pyrénées in France approximately 80 km north-east of the Huesca site revealed the presence of *A. gredana* in France. The species is therefore not endemic to the Iberian Peninsula, though it is likely to have a French range restricted to high altitude sites in the Pyrenees.

Material examined. France: Hautes-Pyrénées, Eget Cité, 4.v.2017, 13, leg. R. Rudelle, R. Rudelle Colln.; Spain: Huesca, San Juan de la Peña, 14.v.1995, 13, leg. H. & J.E. Wiering, RMNH (see also records in Wood et al. 2021).

Subgenus Truncandrena Warncke, 1968

This subgenus contains species that often vary extensively in the colouration of their pubescence, sometimes display minimal variation in structural characteristics in the female sex, and can sometimes only be reliably identified in the male sex. These identification difficulties have led to a large number of subspecific concepts in the literature, the integrity of which must be examined using molecular data. There are a number of taxonomic changes to make which affect the Iberian and more broadly West Mediterranean fauna.

Andrena (Truncandrena) doursana Dufour, 1853 sensu lato

This nominal species is highly variable across its range which was previously considered to be from Morocco and Iberia to Turkey and the Levant. *Andrena doursana* was originally described from Algeria, and Warncke (1975a, 1975b, 1980) described the subspecies *A. d. citreola* Warncke, 1975 from Spain, *A. d. agadira* Warncke, 1980 from southern Morocco, *A. d. bengasia* Warncke, 1980 from Libya, and *A. d. mizorhina* Warncke, 1975 from Turkey. Pisanty et al. (2022b) elevated the eastern subspecies *A. (Truncandrena) doursana mizorhina* to species status, leaving populations in Iberia and north-western Africa to Libya.

The differentiation between these remaining subspecies relies on the colour of the female pubescence, as there are no structural differences in the males; indeed, the subspecies A. d. agadira and A. d. bengasia were described only from the female sex. Andrena d. citreola is bright, and has predominantly white hairs on the face and a light brown terminal fringe with scattered white hairs laterally. Andrena d. agadira is much darker, with dark facial hairs and a uniformly dark terminal fringe. Wood et al. (2020b) found the undescribed female of a similar species endemic to Morocco, A. (Truncandrena) alchata Warncke, 1974, which is structurally very similar to the female of A. d. agadira, but these authors were unable to conclude if the two taxa were synonymous.

Analysis of barcodes from southern and northern Morocco and Iberia shows that female specimens identified as *A. doursana* s. str., *A. d. agadira*, and *A. d. citreola* did not form differentiated clades (Fig. 29). Light forms from southern Portugal [IBIHM1223-22] and the Moroccan Anti-Atlas near Tazenakht [WPATW505-22] were separated, and dark forms from south-western Morocco near Guelmim [WPATW506-22] and Tiznit [WPATW503-22] were clustered closer to the light specimen from Tazenakht. The light specimen from Portugal was clustered close to

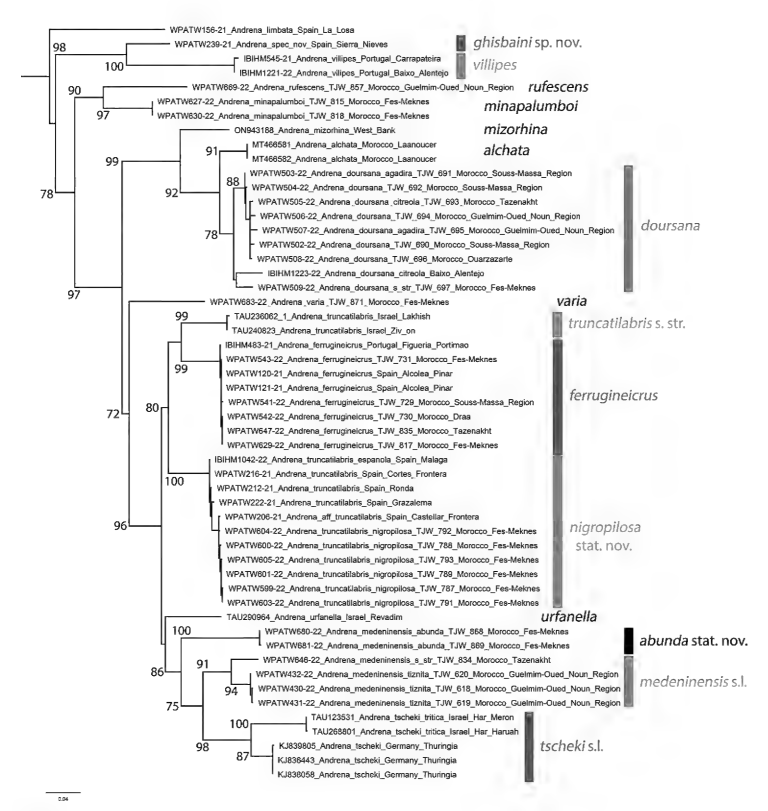


Figure 29. Phylogenetic tree (maximum likelihood) of *Andrena* subgenus *Truncandrena* Warncke, 1968 based on the mitochondrial COI gene. *Andrena* (*Limbandrena*) *limbata* Eversmann, 1852 is used as an outgroup. Numbers adjacent to branches represent bootstrap support (values of <75 are omitted).

a specimen of *A. doursana* s. str. from the Middle Atlas near Taza [WPATW509-22]. This pattern strongly suggests simple separation by distance, with geographically closer specimens displaying more similar barcode sequences. Accepting a broad *A. doursana* species concept, these sequences showed an average intraspecific distance of 2.79% (range 0.29–5.93%).

More broadly, this A. doursana clade had bootstrap support of 78, and was sister to the A. alchata sequences generated by Wood et al. (2020b), a species that has a clearly different male morphology. Andrena alchata has bootstrap support of 91, and was

separated from *A. doursana* by an average of 6.29% (range 5.62–8.03%). Finally, the single sequence of *A. mizorhina* was strongly separated from *A. doursana* by an average of 10.45% (range 10.18–10.94%). These genetic results justify the decision of Pisanty et al. (2022b) to elevate *A. mizorhina* to species level, the findings of Wood et al. (2020b) who identifed the female of *A. alchata*, and the original description of *A. d. agadira* by Warncke as a subspecies, correctly associating dark specimens from south-western Morocco with *A. doursana*. Given the variation in colour forms observed here, no subspecies framework is employed, and so the Iberian taxon is referred to simply as *A. doursana*.

Andrena (Truncandrena) medeninensis Pérez, 1895 sensu lato

Andrena medeninensis was described from Tunisia, and like A. doursana, it nominally displays great variation across its range from Morocco and Iberia to Turkey and the Levant. Warncke (1967, 1974, 1980) described several subspecies, A. m. donata Warncke, 1967 from Spain, A. m. abunda Warncke, 1974 from Morocco, A. m. tiznita Warncke, 1980 from south-western Morocco, and A. m. usura Warncke, 1967 from Turkey. Wood (2023b) synonymised A. m. usura with A. pareklisiae Mavromoustakis, 1957, leaving the North African and Iberian populations.

Sequences of *A. medeninensis* s. str. and *A. m. tiznita* formed a clade with bootstrap support of 91 (Fig. 29). However, the single available sequence of *A. medeninensis* s. str. differed from *A. m. tiznita* sequences by an average of 4.80% (range 4.70–4.85%), *A. m. tiznita* forming a subclade with bootstrap support of 94. However, *A. m. abunda* was strongly separated from this clade of *A. medeninensis* s.l. by an average genetic distance of 10.19% (range 9.97–10.57%). This *A. m. abunda* clade had bootstrap support of 100, and was sister to an *A. medeninensis* s.l. + *A. tscheki* Morawitz, 1872 s.l. clade. *Andrena abunda* stat. nov. is therefore raised to species status; it restricted to Algeria and Morocco. Morphologically, it is extremely similar to *A. medeninensis* s.l., but has much darker pubescence. In the female sex (the male is unknown), the only clear structural difference is that A3 exceeds the length of A4+5+6, whereas in *A. medeninensis* s.l. A3 is shorter than A4+5+6.

As it was unfortunately not possible to sample the Iberian subspecies *A. m. donata*, and no genetic sequences are available from Tunisia, the *locus typicus* for *A. medeninensis* s. str., no further taxonomic action is taken here. Given the large genetic difference displayed by *A. abunda* despite almost no morphological differentiation (at least in the female sex), it is difficult to comment on the Iberian subspecies which simply appears to be a colour variant of this nominally widespread species.

Andrena (Truncandrena) truncatilabris Morawitz, 1877, Andrena (Truncandrena) truncatilabris espanola Warncke, 1967, and Andrena (Truncandrena) truncatilabris nigropilosa Warncke, 1967

Andrena truncatilabris is a widespread species that was originally described from the Caucasus from what is today Armenia (Astafurova et al. 2021). The species is nominally

distributed across the Mediterranean basin, from Morocco and Iberia to the Urals and Iran (Gusenleitner and Schwarz 2002). However, in the east it descends only to the Levant and does not enter the eastern part of North Africa. In the west, Warncke (1967) described two subspecies: *A. t. espanola* from Spain and *A. t. nigropilosa* from Algeria that differed from *A. truncatilabris* s. str. in the structure of their clypeus and their male genital capsule, though Warncke noted that the difference between the two subspecies was minimal, pointing to the darker pubescence of North African specimens as a point of difference.

Sequences from specimens from Spain and Morocco showed almost no genetic differentiation, with an average intraspecific distance of 0.99% (range 0.00–1.85%; Fig. 29). They were strongly separated from *A. truncatilabris* s. str. sequences from northern Israel by an average genetic distance of 9.66% (range 9.12–10.12%). *Andrena truncatilabris* s. str. formed a clade with bootstrap support of 99, and was sister to *A.* (*Truncandrena*) *ferrugineicrus* Dours, 1872, whereas *A. t. nigropilosa* + *A. t. espanola* formed a clade with bootstrap support of 100 that was sister to the *A. truncatilabris* s. str. + *A. ferrugineicrus* clade.

Given this genetic difference, it is clear that specimens from Iberia and north-western Africa are both conspecific and distinct from *A. truncatilabris* s. str. Given this distribution, the use of the name *A. t. espanola* is undesirable, and so *A. nigropilosa* stat. nov. is elevated to species status and *A. t. espanola* syn. nov. is synonymised with it as a subjective junior synonym, as the two names were described in the same publication. The updated synonymy is therefore as follows:

Andrena (Truncandrena) nigropilosa Warncke, 1967, stat. nov.

Andrena (Truncandrena) truncatilabris nigropilosa Warncke, 1967: 225, ♀♂ [Algeria: OÖLM, examined].

Andrena (Truncandrena) truncatilabris espanola Warncke, 1967: 224, ♀♂ [Spain: OÖLM, examined] syn. nov.

Distribution. Portugal, Spain, France, Morocco, Algeria, Tunisia (newly recorded). Material from south-eastern France and northern Italy must be carefully revised, but the position is taken here that the Maritime Alps represent a barrier between *A. nigro-pilosa* and *A. truncatilabris* s. str. This should be confirmed with genetic evidence.

Material examined. ALGERIA: S. Algeria, Laghouat [33.8082°N, 2.8316°E], iii.—iv.1929, 1♀, leg. Meyer, OÖLM (holotype of *A. t. nigropilosa*); Tlemcen, 20.iv.1910, 1♂, leg. de Bergeoin, OÖLM (paratype of *A. t. nigropilosa*); SPAIN: Sierra Nevada [37.0732°N, -3.3948°E], vi.1891, 1♀, leg. Handl., OÖLM (holotype of *A. t. espanola*); Montarco, 28.iv.1924, 1♂, leg. J.M. Dusmet y Alonso, OÖLM (paratype of *A. t. espanola*); Tunisia: Kef, 5 km SW Touiref, 28.iv.2012, 41, leg. C. Sevidy & A. Müller, AMC/TJWC.

Andrena (Truncandrena) villipes Pérez, 1895

Pérez (1895) described A. villipes from north-eastern Spain (Fig. 30), later describing the synonymous A. (Truncandrena) squalida Pérez, 1903 from south-western

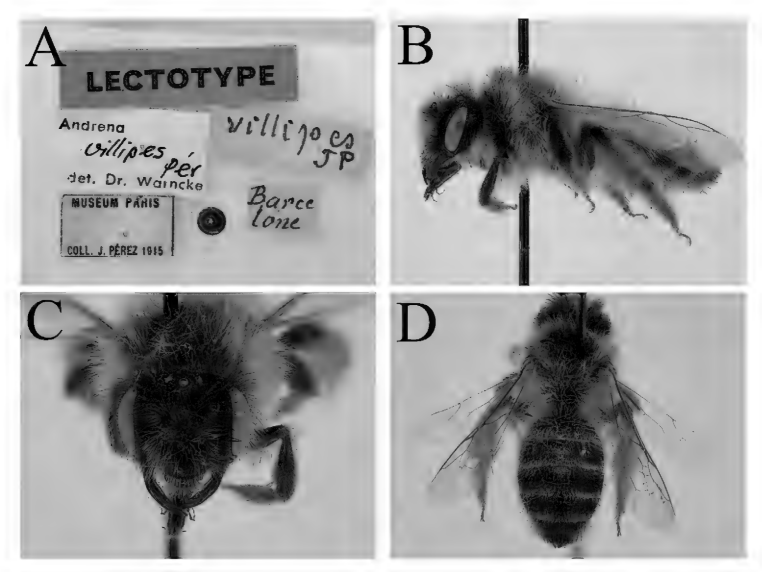


Figure 30. Andrena (Truncandrena) villipes Pérez, 1895; female lectotype **A** label details **B** profile **C** face, frontal view **D** dorsal view.

France. It has a restricted distribution, known from Cistaceae-rich habitats from northern Morocco, Portugal, Spain, southern France, and north-western Italy (Gusenleitner and Schwarz 2002; Lhomme et al. 2020). Searches in the Sierra de las Nieves in southern Spain produced two particularly large and dark specimens provisionally identified as *A. villipes*. A sequence from one of these specimens is separated from *A. villipes* sequences from southern Portugal by an average genetic distance of 11.85% (range 11.85–11.85%). This new species is described and diagnosed below.

Material examined. Spain: Barcelona [41.4028°N, 2.1332°E], 1, MNHN (lectotype of *A. villipes*; Fig. 30).

Undescribed subgenera

As a result of the ground-breaking analysis of Pisanty et al. (2022b), we now have unprecedented phylogenetic resolution for the genus *Andrena*, and are now able to seriously deal with the problems inherent to the subgeneric classification system largely solidified by Warncke (1968a). Building on the work of Pisanty et al. (2022b), the Iberian fauna contains representatives of 44 described subgenera, but also representatives for nine additional clades that are currently undescribed. Some of these are dealt with

here, though the *aegyptiaca*-group (Iberian representative: *A. alluaudi* Benoist, 1961) is not treated here; this lineage will be described in a planned future comprehensive revision of the subgenus.

The former Poliandrena Warncke, 1968

Pisanty et al. (2022b) demonstrated that *Andrena polita* Smith, 1847 falls into an expanded *Ulandrena*, and that *Poliandrena* is a strict synonym of this subgenus, and additionally that *Poliandrena* sensu Warncke contains at least five different polyphyletic clades, with the possibility of additional distinct clades when taxonomic sampling and phylogenetic analysis is more complete. Warncke, who described the subgenus *Poliandrena*, understandably used it as a 'waste-basket' for unclear taxa that displayed no clear defining characters. It is therefore necessary to split up this old grouping and to describe new subgenera.

In Iberia, representatives of all five subgenera can be found. These can be broadly summarised as the blanda-group, the florea-group, the limbata-group, the oviventrisgroup, and the *relata*-group. Four of these five lineages are represented in the analysis of Pisanty et al. (2022b) which is based on UCE analysis. UCE analyses are based on thousands of loci, and hence offer a high degree of confidence that groups are or are not related. Whilst single locus COI analyses cannot compare to those based on UCEs, they can demonstrate if individual species are closely related, and hence complement a UCE-informed phylogeny. No members of the *oviventris*-group were included in the analysis of Pisanty et al. (2022b), but analysis of barcodes (Fig. 31) places A. farinosa Pérez, 1895, A. farinosoides Wood, 2020, and A. oviventris Pérez, 1895 in a single clade with bootstrap support of 99, well-separated from A. corax Warncke, 1975, A. murana Warncke, 1975, and A. relata Warncke, 1975 which belong to the relata-group and which are morphologically the most similar to members of the oviventris-group. The two clades are separated by A. limbata Eversmann, 1852. On the basis of the analysis of Pisanty et al. (2022b) combined with these barcodes, subgenera are described for the blanda-group, florea-group, limbata-group, and oviventris-group.

No action is currently taken for members of the *relata*-group, as the status of morphologically similar species in the Eastern Mediterranean to Central Asia is unclear, and it must be genetically demonstrated if they belong to the *relata*-group or not. The members of the newly described subgenera are detailed below; in an Iberian context, the following species can be considered to be part of the *relata*-group: *A. corax*, *A. laurivora* Warncke, 1974, *A. macroptera* Warncke, 1974, *A. murana*, and *A. relata*.

The caroli-group

Members of this group of species have been placed in the subgenus *Campylogaster* Dours, 1873 (Warncke 1968a) that has the type species *A. erberi* Morawitz, 1871. Pisanty et al. (2022b) identified that *Campylogaster* is polyphyletic, with *A. caroli* Pérez, 1895 falling far away from *A. erberi*. *Campylogaster* therefore can probably only be

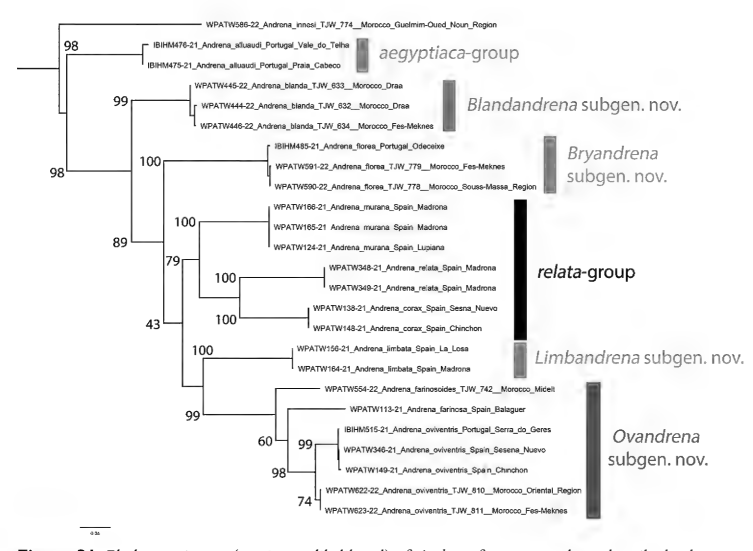


Figure 31. Phylogenetic tree (maximum likelihood) of *Andrena* from currently undescribed subgenera based on the mitochondrial COI gene. *Andrena* (incertae sedis) *innesi* Gribodo, 1894 is used as an outgroup. Numbers adjacent to branches represent bootstrap support (values of <75 are omitted).

applied to the species around *A. erberi* that have strongly depressed tergal margins, e.g. *A. iranella* Popov, 1940 (Iran, Turkemenistan, ?Arabian Peninsula), *A. nanshanica* Popov, 1940 (China, Mongolia), *A. chengtehensis* Yasumatsu, 1935 (China, South Korea). This subgenus therefore appears to be eastern and predominantly Asian, with a western limit of *A. erberi* in the southern Balkans. In contrast, *A. caroli* has the tergal margins flat and not noticeably depressed. Based on the analysis of Pisanty et al. (2022b) combined with this morphological difference, a new subgenus *Pruinosandrena* subgen. nov. is described below for the species around *A. caroli*. Importantly, *A. lateralis* Morawitz, 1876 and *A. incisa* Eversmann, 1852 were also placed into *Campylogaster* by Warncke (1968a). Although not sampled by Pisanty et al. (2022b), *A. lateralis* falls far away from the *Pruinosandrena* based on COI analysis (Fig. 32). Morphologically, *A. lateralis* and *A. incisa* form a species pair (that can be referred to at the *incisa*-group) that lacks many of the characters shared by members of the *Pruinosandrena* (see below), and they probably represent an additional undescribed subgenus. Without additional genetic data, no further steps are taken for this species pair at the present time.

Finally, clarity is required for the status of taxa lumped under a broad concept of *A. pruinosa* Erichson, 1835, specifically *A. pruinosa succinea* Dours, 1872 and *A. pruinosa parata* Warncke, 1967. Erichson (1835) described *A. pruinosa* from southern Spain

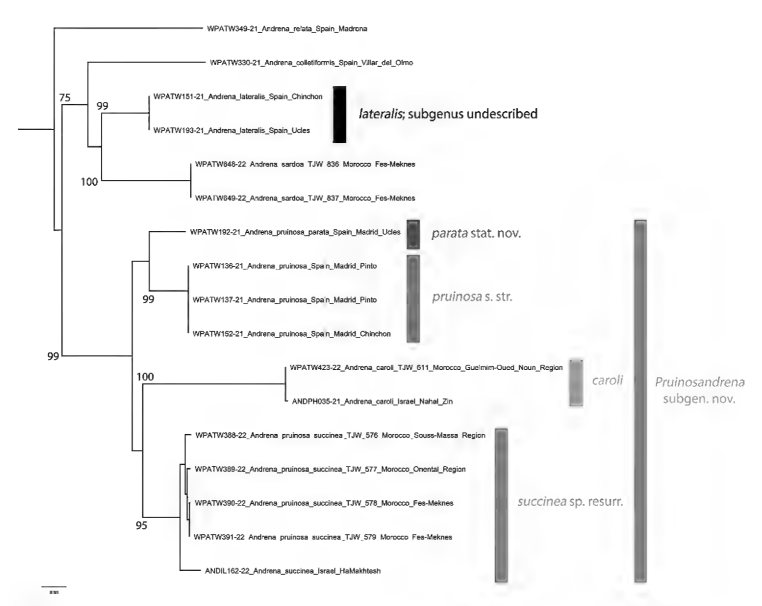


Figure 32. Phylogenetic tree (maximum likelihood) of *Andrena* from the subgenera *Brachyandrena* Pittioni, 1948, *Lepidandrena* Hedicke, 1933, and currently undescribed subgenera based on the mitochondrial COI gene. *Andrena* (incertae sedis) *relata* Warncke, 1975 is used as an outgroup. Numbers adjacent to branches represent bootstrap support (values of <75 are omitted).

(Andalusia); a single female labelled as 'type' is conserved in the ZMHB which may automatically be the holotype (Fig. 33), but this is ambiguous as it is unclear if Erichson described the species from multiple specimens or not. Dours (1872) described *A. succinea* from Algeria, noting the clear red colouration of the metasoma (hence the species name, *succine* amber). Warncke (1967) used *A. succinea* as a subspecies in combination with *A. pruinosa*, arguing that males from North Africa could not be clearly separated from Spanish males. He then described *A. pruinosa parata* from south-eastern Spain, giving characters related to colouration and antennal ratios.

Genetic analysis of members of the *Pruinosandrena* demonstrates that the broad concept of *A. pruinosa* used by Warncke was overly conservative (Fig. 32). A single red-marked female specimen initially identified as *A. pruinosa* was separated by 7.45% from three additional *A. pruinosa* s. str. specimens. Inspection of Warncke's original description of *A. pruinosa parata* and comparison of the antennal ratios demonstrates that this divergent specimen has A3 clearly longer than A4+5, whereas A3=A4+5 in *A. pruinosa* s. str. The other characters mentioned by Warncke relating to colour do not

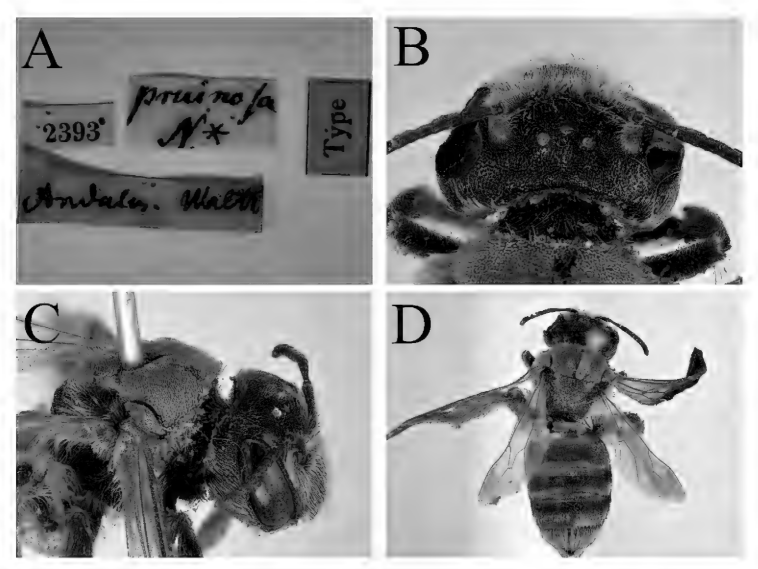


Figure 33. Andrena (Pruinosandrena) pruinosa Erichson, 1835, female holotype/syntype **A** label details **B** head, dorsal view **C** mesosoma, dorsolateral view **D** dorsal view.

work consistently, as one of the barcoded *A. pruinosa* s. str. has the terga partially redmarked and the hind tibiae lightened orange. Though small, this morphological difference is meaningful, as the two taxa are found essentially in direct sympatry; the distance between the sampling location of barcoded specimens of *A. pruinosa* s. str. from Pinto was approximately 11 km from the Camino de Uclés sampling site for *A. pruinosa parata*. In this context, a genetic separation of 7.45% combined with the morphological difference is highly significant, and *A. parata* stat. nov. is treated as a valid species.

Andrena pruinosa succinea was strongly separated from A. pruinosa s. str. by an average genetic distance of 9.45% (range 8.97–10.33%). Sequences of A. pruinosa s. str. were identical, which is not surprising as they all came from a small part of the province of Madrid. Andrena pruinosa succinea samples came from a large geographic area from south-western Morocco to Israel, but still showed low average intraspecific variation of 2.06% (range 0.14–3.80%). The two clades were not sister, being separated by A. parata and A. caroli, and were supported by bootstrap support of 99 and 95, respectively. Andrena succinea sp. resurr. is therefore considered to be a valid species, distinct from A. pruinosa s. str. Morphologically, separation of males is straightforward, and it is unclear why Warncke considered the difference unclear. Andrena succinea males have a yellow marked clypeus (see illustrations in Wood et al. 2020b), with the yellow

markings sometimes extending onto the lower part of the paraocular areas (uniformly black in *A. pruinosa* s. str. and *A. parata*) and, viewed ventrally, A4 is short, as long as broad (A4 is elongate and clearly longer than broad in *A. pruinosa* s. str.). *Andrena parata* males can easily be separated as A3 is longer than A4+5, whereas A3 is shorter than A4+5 in both *A. pruinosa* s. str. and *A. succinea*.

There are also ecological differences. *Andrena succinea* can be found in dry desertedge steppe habitats, as opposed to *A. pruinosa* which in Iberia is found in grasslands and cold steppe that are lightly more lush, at least during the spring. For example, in Morocco, *A. succinea* can be found in stipa steppe habitat around Bou Rached (Oriental region, south of Guercif) on the eastern edge of the Middle Atlas as it transitions into the desert (Fig. 34A, B), whereas it has never been recorded from the more humid parts of the Middle Atlas that have grasslands resembling those that can be found in central Iberia (e.g. Madrid, north of Chinchón, Fig. 34C, D).

Although the type of *A. succinea* is lost, and the type for a more recently described taxon is preserved in the MNHN collection (*A. sitifensis* Pérez, 1895; Fig. 35), *A. succinea* is the name that is established in the literature, either as a species itself or in combination with *A. pruinosa* (Benoist 1961; Warncke 1967; Warncke 1974; Gusenleitner and Schwarz 2002; Wood et al. 2020b; Dermane et al. 2021). In order to conserve this use, a neotype is designated below for *A. succinea*. As a result of these numerous changes, the updated statuses and synonymies are given here:

Andrena (Pruinosandrena) parata Warncke, 1967, stat. nov.

Andrena pruinosa parata Warncke, 1967: 233, ♀♂ [Spain: OÖLM, examined].

Remarks. Though described from south-eastern Spain, Warncke (1976) noted that he had examined a single specimen of *A. parata* (as *A. pruinosa parata*) from Madrid: Ribas de Jarama, but he expressed doubts as to whether or not it had been correctly labelled given the collecting localities of all other known specimens. The contemporary Camino de Uclés site is approximately 7 km from Ribas de Jarama, confirming the presence of this taxon in central Spain.

Distribution. Spain.

Material examined. Spain: Alicante [38.3628°N, -0.5093°W], 1♂, leg. G. Mercet, OÖLM (holotype); Benidorm, 2.vi.1952, 1♀, leg. J. de Beaumont, OÖLM (paratype); Fortuna [Murcia], v.1928, 1♂, leg. J. M. Dusmet y Alonso, OÖLM (paratype); Madrid, Rivas-Vaciamadrid, Canal de Manzanares to Camino de Uclés, 19.v.2021, 1♀, leg. T.J. Wood, TJWC [BOLD accession number WPATW192-21].

Andrena (Pruinosandrena) pruinosa Erichson, 1835

Andrena pruinosa Erichson, 1835: 104, ♀ [Spain: ZMHB, examined].

Andrena lanuginosa Spinola, 1843: 137, ♀ [Spain, lectotype by present designation: MRSN].

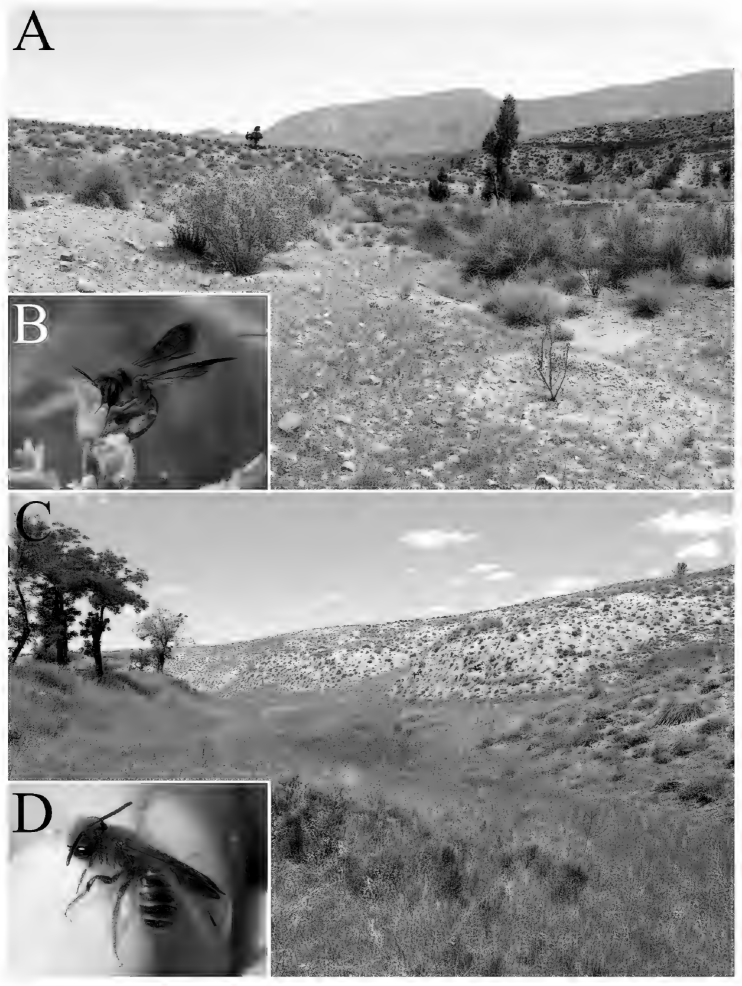


Figure 34. *Andrena* (*Pruinosandrena*) *succinea* Dours, 1872 **A** habitat, Oriental, Guercif, P5427, 2 km SW of Bou Rached, 950 m, 13.v.2022 **B** female collecting pollen from Brassicaceae spp.; *Andrena* (*Pruinosandrena*) *pruinosa* Erichson, 1835 **C** habitat, Madrid, Chinchón, 6 km N, M-311, 14.v.2021 **D** male, in hand.

Remarks. Examination of the type material of both *A. pruinosa* and *A. lanuginosa* (Fig. 36) showed that both were female specimens with dark terga, conforming to the classical concept of this species. Neither represent *A. parata*, and hence the synonymy of *A. lanuginosa* with *A. pruinosa* is maintained. Spinola (1843) did not specify how many specimens he described the species from. The specimen examined here may be automatically the holotype, but as this is not clear from the original description, it is here designated as the lectotype.

Distribution. Spain.

Andrena (Pruinosandrena) succinea Dours, 1872, stat. nov.

Andrena succinea Dours, 1872: 424, ♀ [Algeria: type lost, neotype designated below, OÖLM].

Andrena chrysopyga Dours, 1872: 423, ♀ (nec. Andrena chrysopyga Schenck, 1853) [Algeria: type lost].

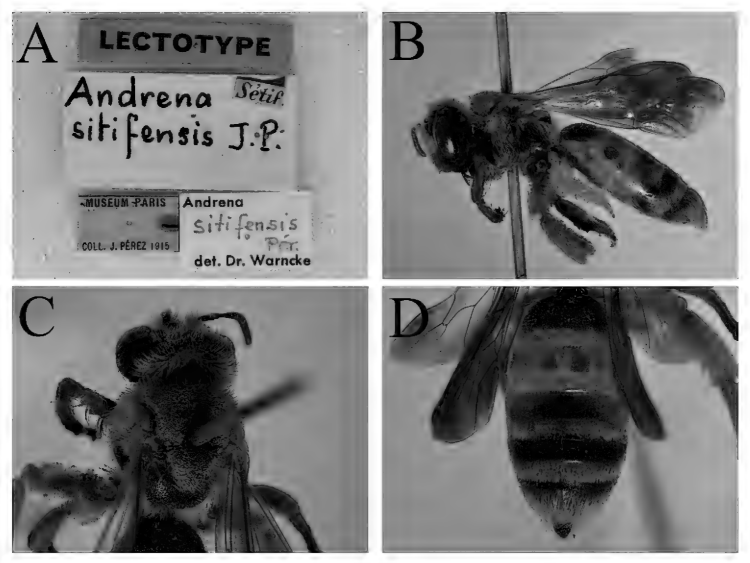


Figure 35. Andrena (Pruinosandrena) sitifensis Pérez, 1895, female lectotype **A** label details **B** profile **C** scutum, dorsal view **D** terga, dorsal view.

Andrena commixta Dalla Torre & Friese, 1895: 43. nom. nov. for Andrena chrysopyga Dours, 1872

Andrena sitifensis Pérez, 1895: 46, ♀ [Algeria: MNHN, examined]
Andrena fulvisquama Popov, 1940: 260, ♀ [Algeria: ZISP, not examined]

Remarks. The synonymy of *A. mayeti* Pérez, 1895 (described from Tunisia) with *A. succinea* reported by Warncke (1967, see also Benoist 1961) is incorrect. Examination of the female lectotype (Fig. 37) shows that *A. mayeti* syn. nov. is a synonym of *A. caroli* Pérez, 1895 (described from Algeria; Fig. 38) because the foveae are wide and there are well-developed hair bands on the tergal margins (see identification key for *Pruinosandrena* below).

Distribution. Morocco, Algeria, Tunisia, Libya, Egypt, Israel and the West Bank, Jordan, Syria, Saudi Arabia, Iran (Wood and Monfared 2022).

Material examined. ALGERIA: Setif [36.2059°N, 5.3965°E], 1♀, MNHN (lectotype of *A. sitifensis*; Fig. 35); Morocco: Oriental, Guercif, P5427, 2 km SW of Bou Rached, 33.8844°N, -3.6154°W, 950 m, 13.v.2022, 1♀, leg. T.J. Wood, OÖLM [BOLD accession number WPATW389-22] (neotype of *A. succinea*, see below).

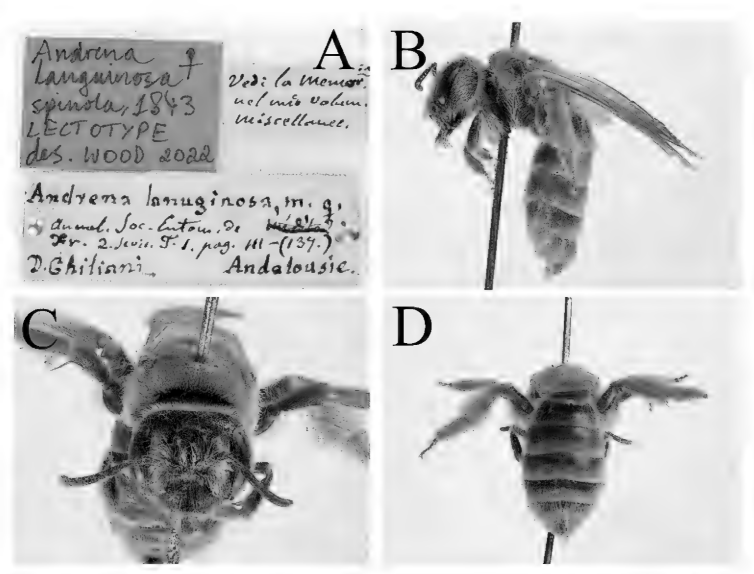


Figure 36. *Andrena* (*Pruinosandrena*) *lanuginosa* Spinola, 1843, female lectotype **A** label details **B** profile **C** head, dorso-frontal view **D** terga, dorsal view.

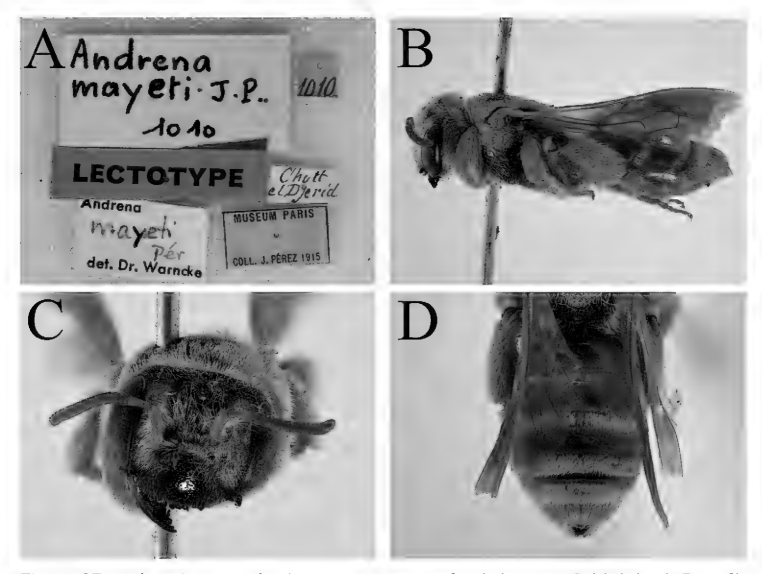


Figure 37. Andrena (Pruinosandrena) mayeti Pérez, 1895, female lectotype **A** label details **B** profile **C** face, frontal view **D** terga, dorsal view.

Andrena (Pruinosandrena) caroli Pérez, 1895

Andrena (Pruinosandrena) caroli Pérez, 1895: 47, ♀ [Algeria: MNHN, examined] Andrena (Pruinosandrena) mayeti Pérez, 1895: 47, ♀ [Tunisia: MNHN, examined] syn. nov.

Distribution. Morocco, Algeria, Tunisia, Egypt, Israel.

Material examined. ALGERIA: Biskra [34.8600°N, 5.6995°E], 1, MNHN (lectotype of *A. caroli*; Fig. 38); **Tunisia:** Chott el Djerid [33.8806°N, 8.1435°E], 1, MNHN (lectotype of *A. mayeti*; Fig. 37).

The numida-group

This group of Palaearctic species was previously placed in the subgenus *Thysandrena* Lanham, 1949 by Warncke (1968a). However, true *Thysandrena* occur only in North America, and Palaearctic species fall elsewhere (Pisanty et al. 2022b). Four species are recognised in the Palaearctic; *A. hypopolia* Schmiedeknecht, 1884, *A. numida*, *A. ranunculorum* Morawitz, 1877, and *A. lunata* Warncke, 1975, though additional

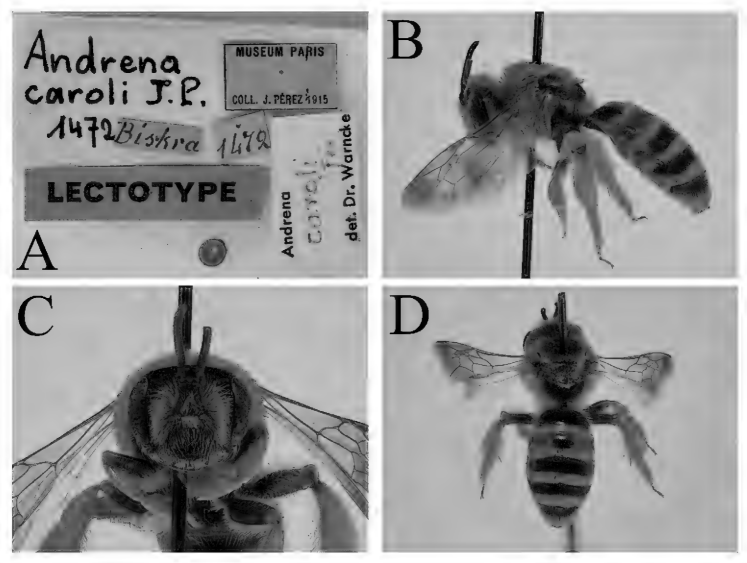


Figure 38. *Andrena (Pruinosandrena) caroli* Pérez, 1895, female lectotype **A** label details **B** profile **C** face, frontal view **D** terga, dorsal view.

genetic work is required to ensure that they all belong together; no subgenus is therefore described in the current work.

The status of *A. hypopolia* (described from southern France) has been somewhat unclear, as no major morphological differences from *A. numida* (described from Algeria) are apparent. Warncke used *A. hypopolia* in combination with *A. numida* as the subspecies for south-western Europe, using several other taxa as subspecies for populations in Central and Eastern Europe (ssp. ?holosericea Bramson, 1879, considered a nomen dubium by Gusenleitner and Schwarz 2002), southern Italy (ssp. syracusae Strand, 1921), and Turkey (ssp. albiscopa Warncke, 1967, see Gusenleitner and Schwarz 2002). No members of this group are known from the Levant.

The situation is further complicated because the type of *A. hypopolia* is lost (and could not be found amongst undesignated Schmiedeknecht type material located in the RMNH collection, see below), and the type of *A. numida* can also not be located at the MNHN. Furthermore, *A. inconspicua* Morawitz, 1871 was described from Calabria in southern Italy. Warncke (1967) treated this as *A. (Taeniandrena) russula* Lepeletier, 1841 based on the description; Gusenleitner and Schwarz (2001) then listed it as a synonym of *A. numida*, before finally Gusenleitner and Schwarz (2002) listed it as a synonym of *A. hypopolia* despite the fact that it is an older name. This is because,

under the broad treatment of Warncke, *A. inconspicua* would be junior to *A. numida*. However, Gusenleitner and Schwarz (2002) treated *A. hypopolia* as a distinct species, and did not resolve the status of *A. inconspicua*, even though it is an older name than *A. hypopolia*. The lectotype of *A. inconspicua* was recently illustrated by Astafurova et al. (2021), confirming its affinity with the *numida*-group, and not the subgenus *Taeniandrena*. The use of the name *A. inconspicua* therefore depends on two things; 1) whether European and North African material are distinct, and 2) whether material from southern Italy shows a stronger affinity with the European or North African taxon.

Genetically, barcoded specimens from Iberia showed almost no differentiation from specimens from Germany (average genetic distance 0.26%; Fig. 1), thus demonstrating that the name A. hypopolia can be applied to both the south-western and Central European populations. However, there was a consistent separation between Moroccan and German/Iberian sequences, these separated by an average genetic distance of 3.05% (range 2.87–3.13%). A single sequence was available from Sicily. This sequence differed from German/Iberian sequences by an average genetic distance of 6.11% (range 3.87-6.82%) and from Moroccan sequences by an average of 2.39% (range 2.17-2.64%). It clustered with Moroccan sequences, with the clades Morocco+Sicily showing bootstrap support of 92 and Germany+Iberia showing bootstrap support of 86. Examination of these barcoded specimens shows that there is a subtle but consistent morphological difference, which is the density of punctures on T3 of females. In Iberian specimens, the disc of T3 is densely punctate, with punctures separated by <1 puncture diameter. In contrast, in Moroccan and Sicilian specimens, the disc of T3 is shallowly and obscurely punctate, with punctures separated by >1 puncture diameter. Examination of the female lectotype of A. inconspicua (Astafurova et al. 2021: fig. 24) shows that this specimen morphologically conforms to A. numida, with weak and obscure punctures on the disc of T3. On this basis, A. inconspicua is synonymised syn. nov. with A. numida, and A. hypopolia is maintained as the senior name for populations in Iberia and Central Europe. A barcoded neotype of A. numida is designated from Moroccan material (see below).

This action largely maintains the *status quo* of Gusenleitner and Schwarz (2002), and *A. numida* is considered to have a distribution of Morocco, Algeria, Tunisia, Libya, and Sicily and the extreme southern parts of Italy (Calabria, Campania). The inclusion of Campania derives from treatment of the taxon *A. syracusae*. This taxon was described by Strand in the male sex only as a variety of *A. (Simandrena) propinqua* Schenck, 1853. Warncke (1967) treated this as a subspecies of *A. numida* with dark facial hair. The barcoded specimen from Sicily is a female, and whilst it should represent *A. syracusae* since it was collected only 150 km west of the *locus typicus* of Syracuse, since the female of *A. syracusae* is unknown and undescribed this cannot be said for certain. I have not seen male *A. numida* material from Sicily, or the type material of *A. syracusae* itself. However, I have seen males from the island of Ischia by Naples. These conform to the concept of *A. syracusae*, with extensive dark facial pubescence, and show extremely weak and sparse punctation on T3. The island of Ischia and the nearby Sorrento Peninsula on the Italian mainland are unusual and host an *Andrena* fauna with

several species that are otherwise found in Italy only in Sicily, such as A. (Micrandrena) fumida Pérez, 1895. On the basis of its morphology, A. syracusae is considered to represent only a form of A. numida. This should ultimately be confirmed through the direct barcoding of males with dark facial pubescence; these are expected to fall into the A. numida clade.

The status of *A. numida albiscopa* is unclear, but based on its distribution and morphology (T3 is clearly punctured), it is transferred to *A. hypopolia albiscopa* comb. nov. The punctures of T3 are sparser than in *A. hypopolia* s. str., and the interspaces are shinier. This is true also of *A. hypopolia* material from Central Asia (Kyrgyzstan). For now, a conservative position is taken that *A. hypopolia* ranges from Iberia to Central Asia and western Siberia, though the eastern limit and the status of material from Turkey requires validation through genetic analysis across this range.

Finally, examination of the lectotype of *A. setosa* Pérez, 1903 (Fig. 39; des. Warncke 1967) reveals that it is not a synonym of *A. numida hypopolia* as given by Warncke (1967), but a synonym (syn. nov.) of *Andrena ranunculorum*. The specimen comes from Arreau in France which is in the Hautes-Pyrénées department, and is situated immediately adjacent to mountains that ascend to 3,000 m. *Andrena ranunculorum* is known from alpine habitats in the central and eastern Pyrenees on both the French and Spanish sides, so this synonymy makes both morphological and ecological sense.

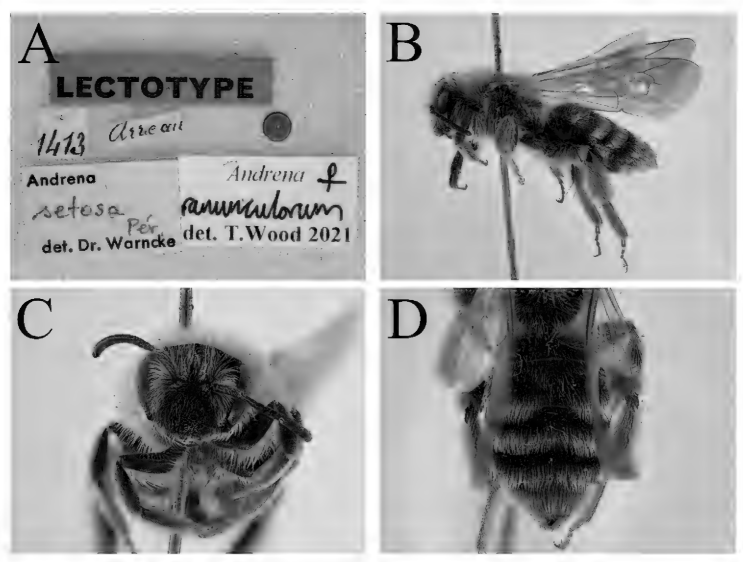


Figure 39. Andrena (incertae sedis) setosa Pérez, 1903, female lectotype A label details B profile C face, frontal view D terga, dorsal view.

Material examined. *Andrena ranunculorum*: France: Arreau [42.9064°N, 0.3557°E], 1♀, MNHN (lectotype; Fig. 39).

Andrena numida f. syracusae: ITALY: Campania, Is. Iscia, Panza, 9.iv.1991, 66, leg. J. Gusenleitner, OÖLM/TJWC.

Andrena fumida: ITALY: Kampanien, Salerno, Monti Alburini (NP), SE Petina, 1100 m, 8.vi.2003, 1♀, leg. H. & R. Rausch, OÖLM; Mondello [Palermo, Sicily], 10.iv.1979, 1♂, leg. J.A.W. Lucas, OÖLM; Monte Faito (Campania), 13.v.1976, 1♂, leg. Pagliano, MRSN; Sorrento [Naples], 3.v.1970, 1♂, leg. J.P. van Lith, RMNH.

Description of new subgenera

Subgenus Blandandrena subgen. nov.

https://zoobank.org/E78F170A-60CC-4D38-9D56-9727B1347DDC

Type species. Andrena blanda Pérez, 1895.

Diagnosis. Blandandrena is monotypic, and hence diagnosis of A. blanda (Fig. 40) is de facto diagnosis of the subgenus. Through the combination of slightly upturned fore margin of the clypeus, fovea broad and occupying over ½ the space between the lateral ocellus and the inner margin of the compound eye, weak but distinct humeral angle, unmodified posterior face of the hind femur (without teeth, carinae, or spines), lack of squamous hairs, simple hind tibial spur (not broadened basally or medially), dark integument, black male clypeus, and essential absence of defining features it falls very close to members of the *relata*-group and to *Ovandrena* subgen. nov. that were formerly lumped together under the subgenus *Poliandrena* (see above). *Andrena blanda* females can provisionally be separated from the relata-group by the weakly punctate terga, punctures shallow and somewhat obscure, separated by 1-2 puncture diameters (Fig. 40D) whereas in the relata-group, tergal punctures are typically much stronger, clear and dense, separated by 1–2 puncture diameters but often by only 1 puncture diameter. This character works for West Mediterranean members of the *relata*-group, but additional work is needed to define this subgenus in the east, and so it is not defined and described here. Andrena blanda females can be separated from the Ovandrena by their simple scutal hairs (Fig. 40A–C; hairs semi-squamous in *Ovandrena*) and poorly delineated propodeal triangle that lacks lateral carinae (Fig. 40C; propodeal triangle clearly defined by presence of lateral carinae in *Ovandrena*; see below for additional detail).

Andrena blanda males can be separated by their black clypeus (Fig. 40F) in combination with their distinctive genital capsule (Fig. 40G). Most members of the relata-group have a yellow-marked clypeus, but for those with a black clypeus (e.g. A. corax), the genital capsule allows recognition. In A. blanda, the genital capsule is short and compact, more or less round, with gonocoxae with apically diverging inner margins and which are produced into short pointed teeth, and the gonostyli are apically strongly broadened and flattened, the disc being slightly broader than long. In the relata-group, the genital capsule is typically simple, without strongly apically broadened gonostyli, or when this is the case (e.g. A. murana Warncke, 1975a) then the clypeus is yellow-marked and the

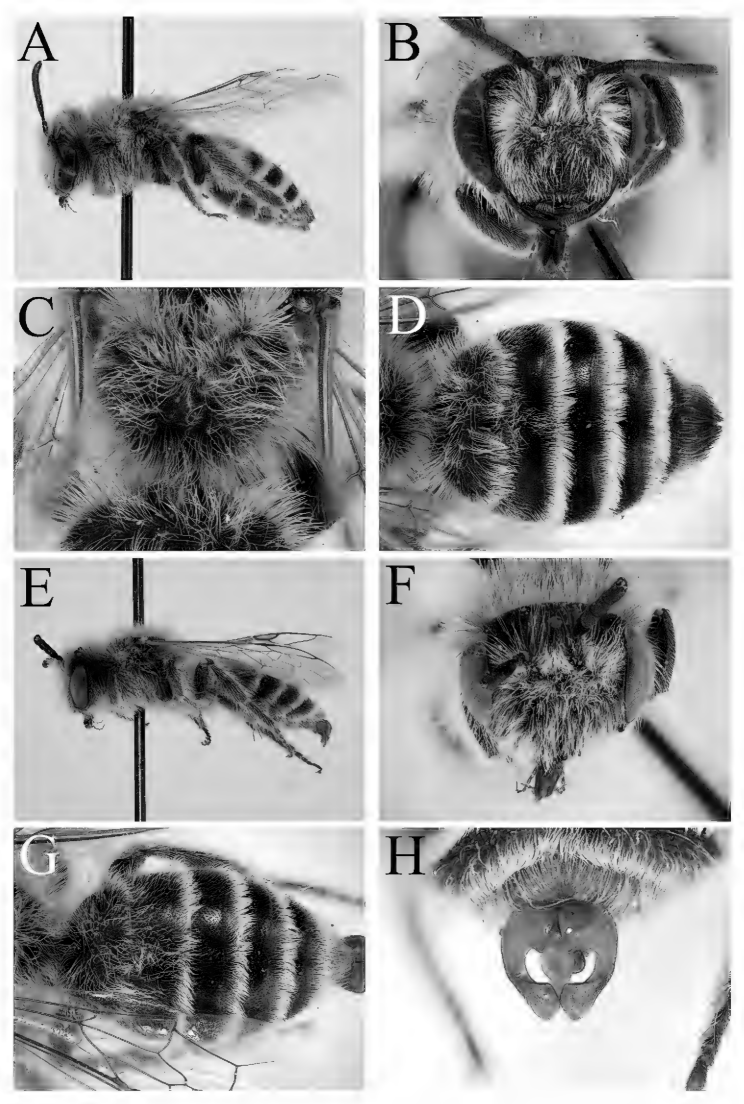


Figure 40. Andrena (Blandandrena) blanda Pérez, 1895, female **A** profile **B** face, frontal view **C** propodeum, dorsal view **D** terga, dorsal view; male **E** profile **F** face, frontal view **G** terga, dorsal view **H** genital capsule.

gonocoxae are not produced into pointed teeth. *Andrena blanda* can be separated from the *Ovandrena* by the genital capsule, as in *Ovandrena* the gonostyli are apically flattened and spatulate, but the disc is always longer than broad. The gonocoxae are also produced into apically projecting teeth, but the inner margins of the gonocoxae are parallel and do not diverge. The propodeal triangle of *Ovandrena* is also triangular and strongly defined by lateral carinae, whereas in *A. blanda* it is poorly defined and lacks lateral carinae.

Description. Medium-sized bees (9–10 mm) with dark integument. Head broad, 1.4 times broader than long. Gena slightly exceeding width of compound eye; ocelloccipital distance 1.5–2 times diameter of lateral ocellus, slightly broader in male sex. Facial fovea broad, occupying almost entire distance between lateral ocellus and inner margin of compound eye. Mesosoma dorsally with moderately long light brown hairs, laterally with white hairs. Pronotum laterally with humeral angle. Dorsolateral surface of propodeum with obscure and finely raised rugosity; propodeal triangle broad, poorly delineated laterally, surface with fine granular reticulation, basally with weakly raised rugosity, propodeal triangle thus defined by change in surface sculpture. Forewing with nervulus antefurcal. Hind tibial spurs simple, not broadened basally or medially. Terga weakly and obscurely punctate, punctures separated by 1–2 puncture diameters. Male genital capsule rounded, more or less circular in outline, gonocoxae with inner margins apically diverging, produced into apically projecting short pointed teeth. Gonostyli apically broadened and flattened, apical disc slightly broader than long. Penis valves relatively narrow, occupying less than half space between gonostyli.

Etymology. The name is taken from the name of the type species *A. blanda*, with *blanda* being the feminine singular of the adjective *blandus* which can mean pleasant, agreeable, smooth. It can be used to refer to the generally unremarkable nature of the species which has made it hard to assign to a particular group of species. The gender is feminine.

Included species. *Andrena blanda* (Spain, including mainland Spain and newly recorded for Fuerteventura), Morocco, Algeria, Tunisia; Gusenleitner and Schwarz 2002).

Material examined (illustrative). Algeria: Biskra [34.8600°N, 5.6995°E], 1♀, MNHN (lectotype des. Warncke 1967); 5 km N of Mecheria, st. 9, 7.iv.1983, 1♀, leg. R. Leys & P. v. d. Hurk, RMNH; 5 km SE of Sfissifa, st. 8, 6.iv.1983, 1♀, leg. R. Leys & P. v. d. Hurk, RMNH; Morocco: Drâa-Tafilalet, Tazenakht, 1 km W Anezal, c. 1600 m, 15.iv.2022, 1♀, leg. T.J. Wood, TJWC; Foum Zguid, 50 km N, 30.iii.1986, 1♂, 16♀, leg. M. Schwarz, MSC; Ifkern, 25 km E Boulemane, 25.v.1995, 10♀, leg. Mi. Halada, OÖLM; Spain: Fuerteventura, Costa Calma, ESE Montaña Pelada, 29.iii.2015, 3♀, leg. A. Müller, AMC/TJWC; Fuerteventura, S Costa Calma, Montañeta de los Verdes, 1.iv.2015, 5♀, leg. A. Müller, AMC; Sierra de María, 25 km W Lorca, 10.v.2003, 1♀, leg. J. Halada, OÖLM; Granada, Pantano de los Bermejales, 26.v.1982, 1♀, leg. R. Leys, RMNH.

Subgenus Bryandrena subgen. nov.

https://zoobank.org/54B1C9C7-8D5A-441B-B49B-DDBA8FB5FE39

Type species. Andrena florea Fabricius, 1793.

Diagnosis. Bryandrena is monotypic, and hence diagnosis of A. florea is de facto diagnosis of the subgenus. The combination of broad head (Fig. 41B, F; 1.3–1.4 times broader than long), the inner margins of the compound eyes diverging ventrally, humeral angle (stronger in male), punctate clypeus with underlying surface shiny over the majority of its area, propodeal triangle not defined by lateral carinae and comparatively smooth relative to the microreticulate dorsolateral parts of the propodeum (Fig. 41C), terga that are always at least partially red-marked (Fig. 41D, G), unmodified posterior face of the hind femur (without teeth, carinae, or spines), lack of squamous hairs, simple hind tibial spur (not broadened basally or medially), black male clypeus, and unique genital capsule (Fig. 41H; see description below) allows separation from any other Andrena species.

Description. Medium-sized bees (11–13 mm), integument predominantly dark, with red markings on at least one tergum, sometimes all terga extensively red marked. Head broad, 1.3–1.4 times wider than long, inner margins of compound eyes diverging ventrally. Gena slightly exceeding width of compound eye in females, clearly exceeding width of compound eye in males; ocelloccipital distance 1.5–2 times diameter of lateral ocellus. Facial fovea broad, occupying 3/4 of distance between lateral ocellus and inner margin of compound eye. Pronotum laterally with humeral angle, more pronounced in male sex. Dorsolateral surface of propodeum microreticulate, with weakly raised reticulation; propodeal triangle poorly defined laterally, comparatively smooth and lacking microreticulation, basally with raised rugosity covering variable extent, never entire propodeal triangle. Forewing with nervulus interstitial. Terga regularly and densely punctate, punctures separated by 1 puncture diameter. Male genital capsule strongly elongate, gonocoxae essentially truncate with inner margin rounded, gonostyli apically produced, elongate, strongly flattened and spatulate, 3 times longer than broad; penis valves basally broad, strongly narrowing medially to become elongate and acutely pointed apically.

Etymology. The name is taken from the pollen host plant *Bryonia* (Cucurbitaceae) which ultimately derives from the Greek βρυωνία [bruōnía]. *Andrena florea* can be found frequently almost wherever *Bryonia* species are in flower. The gender is feminine.

Included species. Andrena florea (West Palaearctic, from Morocco and Iberia to Iran and the Ural Mountains; Gusenleitner and Schwarz 2002).

Subgenus Limbandrena subgen. nov.

https://zoobank.org/E0948E05-DB8C-4AD2-B260-01D335F6EA96

Type species. Andrena limbata Eversmann, 1852 (illustrated by Astafurova et al. 2022).

Remarks. Historically, *A. toelgiana* Friese, 1921 has been considered the sister species to *A. limbata*, differing by the yellow clypeus in the female sex (see Gusenleitner and Schwarz 2002). However, the male is unknown. Structurally, there are no differences. Examination of specimens from Bulgaria and Turkey show a gradient of yellow colouration on the clypeus, from entirely black, with a narrow longitudinal yellow

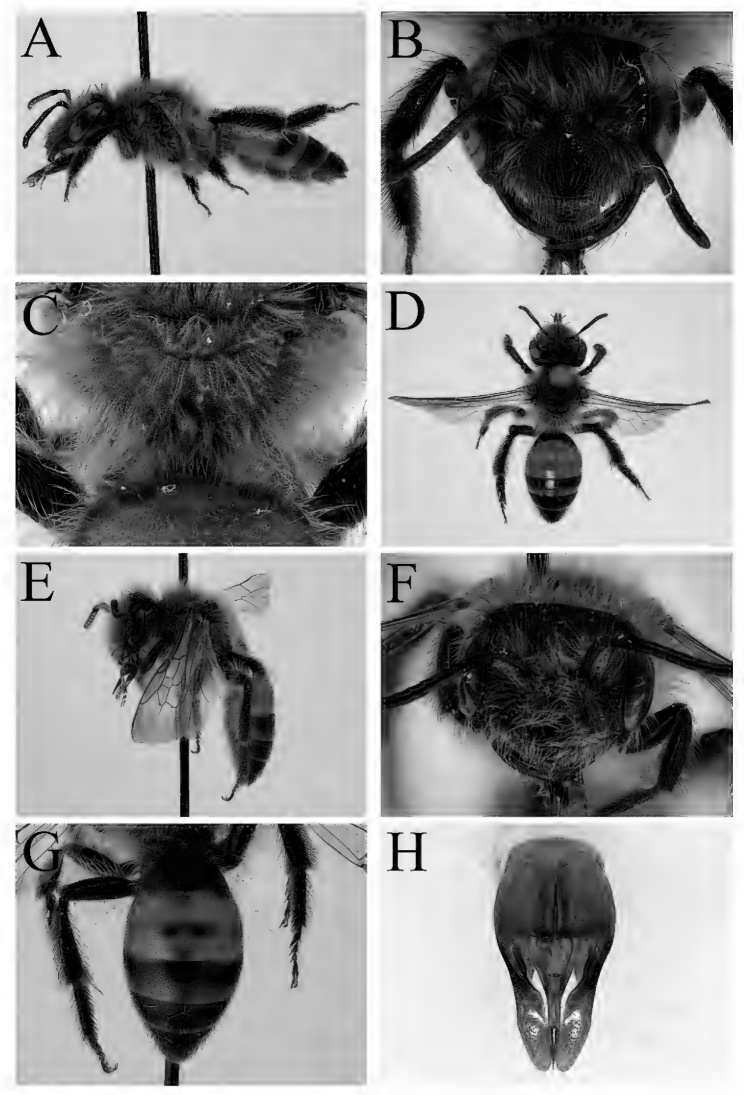


Figure 41. Andrena (Bryandrena) florea Fabricius, 1793, female **A** profile **B** head, frontal view **C** propodeum, dorsal view **D** dorsal view; male **E** profile **F** face, frontal view **G** terga, dorsal view **H** genital capsule.

strip, predominantly yellow-marked, and entirely yellow-marked (Fig. 42). Across this gradient there is also variation in the strength of the tergal hair bands, with material nominally conforming to *A. toelgiana* falling closer to *A. limbata* s. str. in displaying clear hair bands, whereas the subspecies *A. limbata dusmeti* Warncke, 1975 (Portugal, Spain, southern France, north-western Italy) has hair bands that are almost absent in fresh specimens (Figs 43C–H, 44A, B, E, F). On the basis of this colour gradient, absence of structural characters, and lack of a described male that could offer a distinctive difference, *A. toelgiana* syn. nov. is synonymised with *A. limbata*.

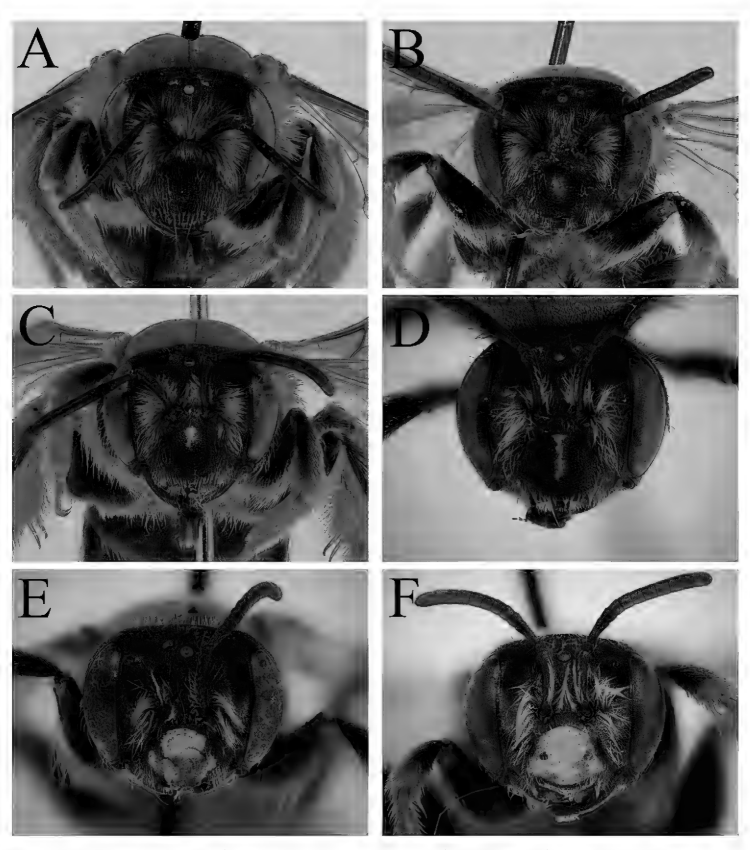


Figure 42. Andrena (Limbandrena) limbata Eversmann, 1852 female head in frontal view on west-east gradient **A** Spain **B** Bulgaria **C** Turkey (Gürün) **D** Turkey (Bingöl) **E** Turkey (Bingöl) **F** Turkey (Diyarbakır).

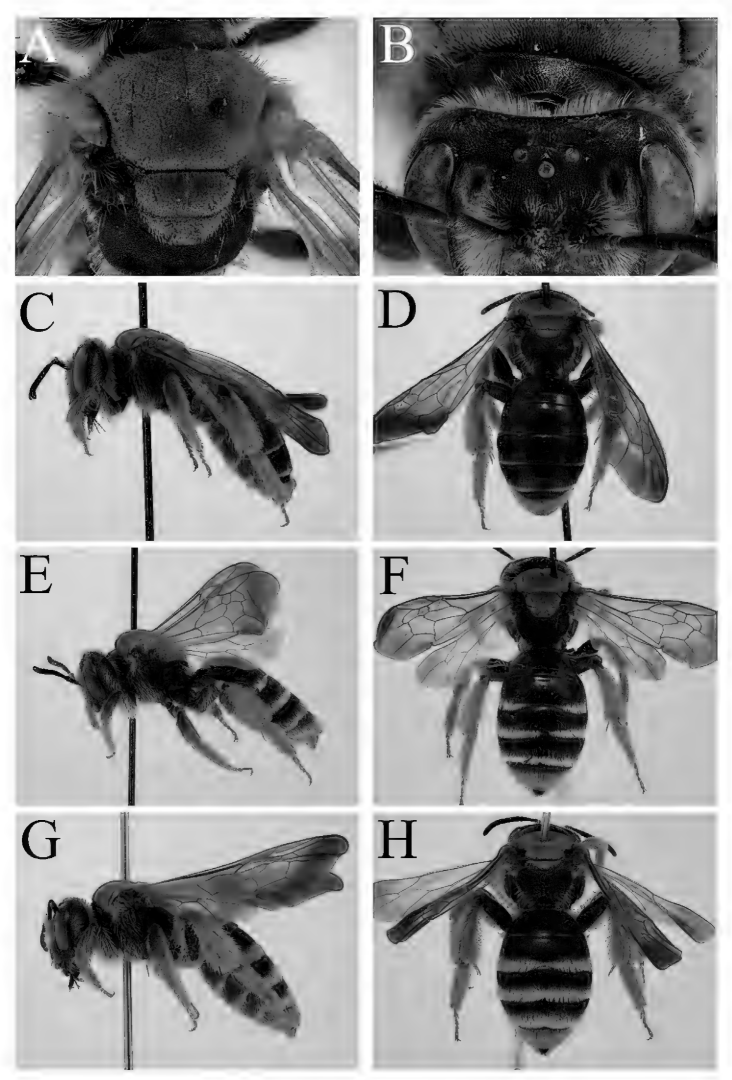


Figure 43. Andrena (Limbandrena) limbata Eversmann, 1852, female **A** scutum, dorsal view **B** head, dorso-frontal view; Andrena limbata dusmeti Warncke, 1967 female **C** profile **D** terga, dorsal view; A. limbata s. str. female, Bulgarian specimen **E** profile **F** terga, dorsal view; A. limbata s. str. female, Turkish specimen **G** profile **H** terga, dorsal view.

Diagnosis. Limbandrena (and, de facto, A. limbata) can be recognised in the female sex due to the combination of squamous brown hairs on the scutum, scutellum, and metanotum (Fig. 43A), the long ocelloccipital distance equal to three times the diameter of a lateral ocellus (Fig. 43B), the posterior face of the hind femur which lacks a transverse carina and transverse row of raised teeth or spines, the weakly and shallowly punctate dorsolateral surfaces of the propodeum, the laterally clearly delineated and internally finely rugose propodeal triangle (not rugosely areolate), the pronotum with at most a weak lateral angle, the more or less squarish head which is only 1.1 times wider than long, and the simple hind tibial spur that is not broadened basally or medially.

Males can be recognised by most of the same characters: the yellow clypeus (Fig. 44C, D), the relatively rounded head which is only 1.1–1.2 times broader than long (Fig. 44C, D), the long ocelloccipital distance equal to three times the diameter of a lateral ocellus, the pronotum with at most a weak lateral angle, the weakly and shallowly punctate dorsolateral surfaces of the propodeum, the laterally clearly delineated and internally finely rugose propodeal triangle (not rugosely areolate), and the genital capsule with weakly produced rounded gonocoxal teeth, gonostyli with weakly raised and rounded projection on inner margin, and penis valves with rounded lateral hyaline extensions (Fig. 44G, H). No other *Andrena* species shows this combination of characters.

Description. Medium-sized bees (11–14 mm) with dark integument with exception of yellow maculations on female (sometimes) and male clypeus (always). Head 1.1–1.2 times broader than long, compound eyes with inner margins weakly converging apically. Gena slightly exceeding width of compound eye; ocelloccipital distance long, 3 times diameter of lateral ocellus. Facial fovea moderate, occupying ½ space between lateral ocellus and compound eye. Female scutum, scutellum, and metanotum covered with short brown squamous hairs. Pronotum laterally with weak humeral angle. Dorsolateral surface of propodeum weakly and shallowly but regularly punctate, punctures separated by 0.5–1 puncture diameter. Propodeal triangle clearly delineated laterally by raised carinae, internal surface with clear pattern of fine rugosity medially, not extending over entire area. Forewing with nervulus strongly postfurcal. Hind tibial spurs simple, not broadened basally or medially, apically weakly bent. Terga densely and finely punctate, punctures separated by 0.5 puncture diameters. Male genital capsule slightly elongate, with produced and weakly rounded gonocoxal teeth. Gonostyli with weakly raised and rounded projection on inner margin. Penis valves produced into rounded hyaline extensions laterally, occupying majority of space between gonostyli.

Etymology. The name is taken from the name of the type species *A. limbata*, with *limbata* being the feminine singular of the adjective *limbatus* which means edged or fringed, probably in reference to the distinct squamous hairs on the female scutum and scutellum. The gender is feminine.

Included species. *Andrena limbata* (Europe from Portugal and Spain to Turkey, Israel, northern Iran, and the Ural Mountains; Gusenleitner and Schwarz 2002).

Material examined (illustrative). Albania: Lopan [Lapanj], 14.vi.2018, 2♀, leg. Kobe Janssen collection (Belgium); Bulgaria: Lozenec [Lozenets, Λозенец]/Mičurin, 24.vi.1988, 5♀, leg. В. & O. Tkalců, OÖLM; Croatia: Istrien, Rovinjsko Selo, 8–9.

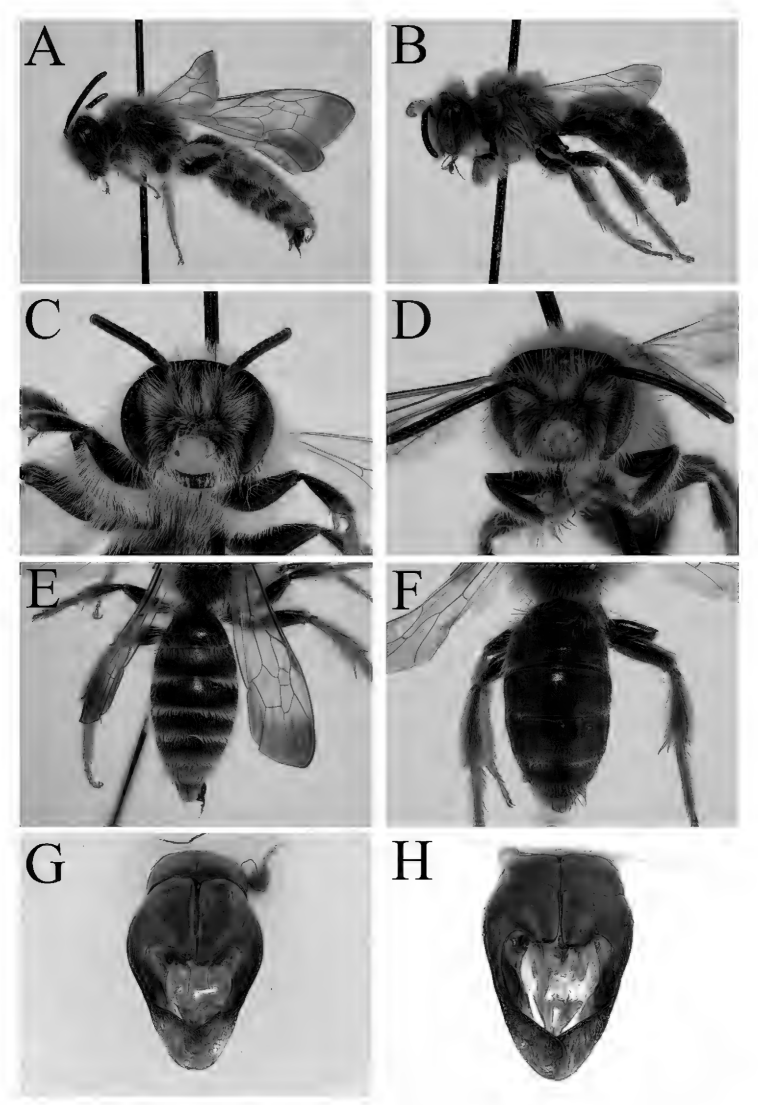


Figure 44. Andrena (Limbandrena) limbata Eversmann, 1852, Turkish male **A** profile **C** face, frontal view **E** terga, dorsal view **G** genital capsule; Andrena limbata dusmeti Warncke, 1967, Spanish male **B** profile **D** face, frontal view **F** terga, dorsal view **H** genital capsule.

vi.2012, $1\cappe$, leg. Holzmann, OÖLM; France: B. d. R., Fontvieille, 28.v.1993, $4\cappe$, leg. H. & J.E. Wiering, RMNH; Georgia: E Tbilisi, lori river S Sagarejo, 19.vi.2015, $3\cappe$, leg. M. Snižek, OÖLM; Greece: Kos I. Kefalos, 4.v.1989, $2\cappe$, leg. K.M. Guichard, NHMUK; Peloponnese, 12 km NWW Sparti, 26.v.2005, $1\cappe$, leg. M. Kadlecová, OÖLM; Italy: Arezzo, Oliveto, 28.v.1989, $3\cappe$, leg. H. & J.E. Wiering, RMNH; Piemonte, S. Benefetto Belbo, 10.vi.1979, $1\cappe$, leg. Pagliano, MRSN; Portugal: Manteigas, Serra da Estrela [40.4009°N, -7.5396°E], 28.iv.-9.vii.1929, $1\cappe$, leg. Kricheldorf, OÖLM (holotype of *A. limbata dusmeti*); Spain: Segovia, Madrona, 500 m NE, Arroyo del Hocino, 15.v.2021, $1\cappe$, leg. T.J. Wood, TJWC; Ávila, Hoyocasero, 1350 m, 20.v.1995, $1\cappe$, leg. H. & J.E. Wiering, RMNH; Turkey: Tanin-Tanin-Pass, 1700 m, 12.vi.1984, $1\cappe$, OÖLM; Karatepe/Adana, 17.iv.1984, $3\cappe$, leg. K. Warncke, OÖLM; Akyaka, Yeşilova, 20.vi.2016, $1\cappe$, leg. M. Kasparek, OÖLM.

Subgenus Ovandrena subgen. nov.

https://zoobank.org/A122C278-C33D-4790-A34A-FC270AE6A962

Type species. Andrena oviventris Pérez, 1895.

Diagnosis. Through the combination of slightly upturned fore margin of the clypeus, broad fovea occupying at least ½ the space between the lateral ocellus and the inner margin of the compound eye, weak but distinct humeral angle, unmodified posterior face of the hind femur (without teeth, carinae, or spines), simple hind tibial spur (not broadened basally or medially), dark integument, black male clypeus, and essential absence of defining features it falls very close to members of the *relata-*group and to *Blandandrena* that were formerly lumped together under the subgenus *Polian-drena* (see above).

In the female sex, Ovandrena species differ from these groups by only a single major character: the structure of the propodeal triangle (Fig. 45A, B). Ovandrena species have scutal hairs that are not as short and clearly squamous as in members of the Aenandrena, Limbandrena, or Lepidandrena Hedicke, 1933, but they are partially squamous in some species (Fig. 45C, D), whereas the scutal hairs in the relata-group and Blandandrena are typically simple, though in some species such as A. corax the hairs can be semi-squamous. The structure of the propodeal triangle is therefore the only character that allows consistent separation. In *Ovandrena*, the propodeal triangle is a broad and well-defined triangle (lateral margins extending almost to the lateral edges of the metanotum) that is defined by raised carinae; the internal surface is covered in raised irregular carinae of a similar width, therefore appearing homogenous and consistent (Fig. 45A, B). In the *relata*-group and *Blandandrena*, the propodeal triangle varies from smooth and undefined to weakly defined with at most irregular rugosity on its internal surface, but never approaches the condition in *Ovandrena*. The combination of these two characters therefore always allows recognition. They can also be partially recognised by the extremely dense and fine punctation of the terga, punctures separated by <0.5 puncture diameters; in comparison groups, the punctures are never this fine or this dense.

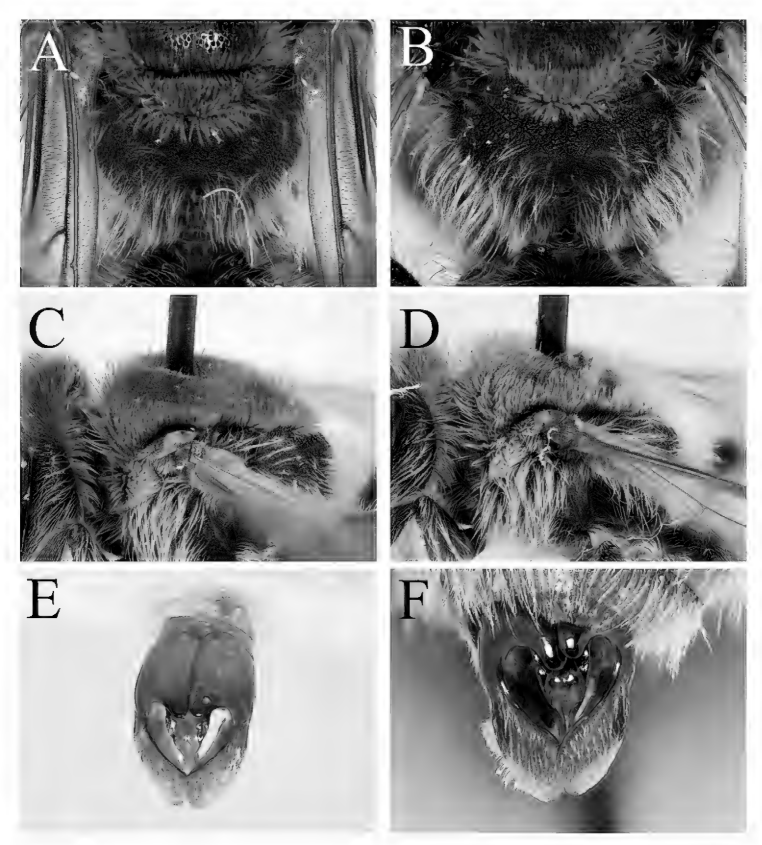


Figure 45. Andrena (Ovandrena) subgen. nov. characters. Andrena (Ovandrena) marsae Schmiedeknecht, 1900, female **A** propodeal triangle; Andrena (Ovandrena) oviventris Pérez, 1895 **B** female propodeal triangle **C** female scutal hairs, profile view **F** male genital capsule; Andrena (Ovandrena) farinosa Pérez, 1895 **D** female scutal hairs, profile view **E** male genital capsule.

Males can be recognised through their combination of dark clypeus with upturned fore margin, distinctive propodeal triangle as in the female sex, pronotum with weak or strong humeral angle, A3 exceeding A4 but shorter than A4+5, and genital capsule which is compact with pronounced and rounded gonocoxal teeth (Fig. 45E, F). Members of the *relata*-group often have a yellow-marked clypeus, but for species showing a dark clypeus (e.g. *A. corax*), the clearly defined triangular propodeal triangle allows separation (smooth to at most weakly defined in the *relata*-group). Separation

can be made from *Blandandrena* by the genital capsule and the propodeal triangle, as detailed in the diagnosis section for that subgenus.

Description. Small to moderately sized bees (7–11 mm) typically with dark integument, one species with red tergal markings; male clypeus always dark. Head broad, 1.3–1.4 times broader than long, compound eyes with inner margins weakly converging apically. Gena slightly exceeding width of compound eye; ocelloccipital distance short, slightly less than to slightly more than diameter of lateral ocellus. Facial fovea moderately broad, occupying ½ distance between lateral ocellus and compound eye. Female scutum, scutellum, and metanotum covered with shortish light brown to whitish semi-squamous hairs (Fig. 45C, D). Pronotum laterally with weak to strong humeral angle. Dorsolateral surface of propodeum with regular and dense network of raised reticulation; propodeal triangle wide, lateral margins extending almost to lateral edges of metanotum, laterally defined by straight raised carinae, thus strongly triangular (Fig. 45A, B). Internal surface of propodeum covered by raised irregular carinae of a similar width, therefore homogenous and consistent. Forewing with nervulus interstitial to weakly antefurcal. Hind tibial spurs simple, not broadened basally or medially. Terga densely punctate, in females punctures extremely dense and fine, separated by < 0.5 puncture diameters, in males punctures slightly sparser and coarser, separated by 0.5-1 puncture diameters. Male genital capsule compact, consistent across species, with gonocoxae strongly produced into apical rounded teeth (Fig. 45E, F). Gonostyli apically spatulate, with a more or less strongly raised inner margin. Penis valves basally narrow or moderately broad, in both cases becoming narrow medially.

Etymology. The name is taken from the type species for the genus, *A. oviventris*. It derives from the Latin *ovum* meaning egg, in reference to the egg-shaped metasoma. The gender is feminine.

Included species. Andrena farinosa Pérez, 1895 (Spain and France), Andrena farinosoides Wood, 2020 (Morocco), Andrena marsae Schmiedeknecht, 1900 (Morocco, Algeria, Tunisia), and Andrena oviventris (Morocco, Algeria, Portugal, Spain, France). The subgenus is therefore currently restricted to the Western Mediterranean, and the centre of diversity is Morocco. The status of A. (incertae sedis) inusitata Pisanty, 2022 must be resolved through genetic analysis (see Pisanty et al. 2022a).

Material examined. *Andrena oviventris*: ALGERIA: Teniet el Had [35.8727°N, 2.0007°E], 1♀, MNHN (lectotype; Fig. 46).

Andrena farinosa: Spain: Murcie [Murcia], 12, MNHN (lectotype).

Identification key to subgenus Ovandrena subgen. nov.

1	Females	. 2
_	Males ¹	.5

¹ Note, males of these four species are extremely similar, and great care must be taken when separating them. Association with females should be made, as well as reference to their known biogeographical distributions and to confidently determined reference material.

2	Terga almost entirely red-marked, with at most slight black marks basally on T1 and two black spots laterally on T2 (north-western Africa)
- 3	Terga dark, without red markings
_	Scutum densely punctate, punctures separated by 0.5 puncture diameters to confluent, narrow interspaces shiny
4	Tergal discs glabrous, with hairs restricted to marginal areas. Larger, 10–11 mm (north-western Africa and south-western Europe)
-	Tergal discs extensively covered with extremely short hairs, forming a velvety pubescence in addition to denser and longer hairs on tergal margins. Smaller, 8–9 mm (Spain and France only)
5	Larger, 9–10 mm. Tongue with outer surface of galea clearly punctate, punctures separated by 1–2 puncture diameters. Sterna with weak and sparse fringes on apical margins. Tergal punctation comparatively larger and coarser (north-western Africa and south-western Europe)
_	Smaller, 7–8 mm. Tongue with outer surface of galea more or less smooth and shiny, without obvious punctures. Sterna with strong and dense fringes on apical margins. Tergal punctation comparatively fine
6	Clypeus comparatively less densely punctate, punctures separated by 0.5–1 puncture diameters, with shiny interspaces, thus appearing shiny. Pronotum with humeral angle comparatively strong (Spain and France only)
_	Clypeus comparatively more densely punctate, punctures separated by 0.5 puncture diameters, interspaces dull, thus appearing dull (north-western Africa)
7	rica)
-	Tergal and sternal margins dark to hyaline, never extensively lightened red- dish orange-brown (Morocco only) ²

Subgenus Pruinosandrena subgen. nov.

https://zoobank.org/62A1E8AF-01AD-4D69-8CD5-E20843A286D5

Type species. Andrena pruinosa Erichson, 1835.

² These two species are almost identical in the male sex, and no structural characters are currently apparent.



Figure 46. Andrena (Ovandrena) oviventris Pérez, 1895, female lectotype **A** label details **B** profile **C** face, frontal view **D** terga, dorsal view.

Diagnosis. This group of species was formerly placed in the subgenus Campylogaster due to the unusual character of the strongly and densely punctate mesepisternum and dorsolateral parts of the propodeum (punctures separated by <0.5 puncture diameters). However, as discussed above, Campylogaster sensu Warncke is polyphyletic and can be broken into three distinct clades (Campylogaster s. str., *Pruinosandrena*, and the *incisa*-group). All three share the distinctly punctate mesepisternum and also an extremely long ocelloccipital distance exceeding three times the diameter of the lateral ocellus. However, separation is straightforward. The true Campylogaster have the marginal area of the terga clearly and distinctly impressed with the apical margin reflexed; the impression therefore forms a latitudinal depressed furrow. In *Pruinosandrena*, the marginal areas of the terga are flat, without any kind of depression. In the *incisa*-group, the tergal margins are flat and the mesepisternum is densely punctate, but the dorsolateral parts of the propodeum have only raised reticulation, without punctures. The propodeal triangle is also clearly delineated by raised lateral carinae, whereas these are absent in *Pruinosandrena*. The combination of extremely long ocelloccipital distance, dense punctures on the mesepisternum and dorsolateral parts of the propodeum, and flat tergal marginal areas is therefore unique and characterises Pruinosandrena.

Description. Medium-sized bees (10-14 mm). Integument variable, from dark with at most tergal margins lightened hyaline-yellow to entirety of metasoma and legs red-marked; male clypeus yellow-marked in one species. Head moderately broad, 1.2 times broader than long. Gena broad, exceeding width of compound eye; ocelloccipital distance extremely long, at least 3 times diameter of lateral ocellus. Facial fovea variable, from narrow to occupying entirety of distance between lateral ocellus and inner margin of compound eye. Female scutum, scutellum, and metanotum with pubescence variable, in some species with extremely short squamous hairs, hairs longer and non-squamous in other species. Pronotum laterally with humeral angle. Mesepisternum and dorsolateral parts of propodeum densely and clearly punctate, punctures confluent to separated by <0.5 puncture diameters. Propodeal triangle without lateral carinae, internal surface with dense network of irregularly raised rugosity, thus contrasting punctate dorsolateral surface. Forewing with nervulus interstitial. Hind tibial spurs simple, not broadened basally or medially. Terga typically densely and finely punctate, punctures separated by 1 puncture diameter, at least on T2-5. Male genital capsule simple, compact, with gonocoxae apically truncate to produced into weak rounded teeth. Gonostyli apically spatulate, penis valves more or less narrow, parallel-sided, occupying less than ½ space between gonostyli.

Etymology. The name is taken from the type species for the subgenus, *A. pruinosa*. The Latin word *pruinosa* is the feminine singular of *pruinosus* which means 'frosty', in reference to the squamous hairs of the mesosoma. The gender is feminine.

Included species. Andrena caroli (Morocco to Israel); Andrena nilotica Warncke, 1967 (Spain); Andrena parata (Spain); Andrena pruinosa (Spain); Andrena sparsipunctata Wood, 2020 (Morocco); Andrena succinea (Morocco to Iran and Saudi Arabia). The centre of diversity is therefore Spain + Morocco, with all six species occurring here.

Identification key to subgenus Pruinosandrena subgen. nov.

Note, the male of *A. sparsipunctata* is unknown. The males described by Wood et al. (2020b) are actually misidentified *A. mediovittata* Pérez, 1895; *A. sparsipunctata* is expected to be restricted to south-western Morocco. See Wood et al. (2020b) for additional useful images.

1	Females2
_	Males
2	In dorsal view, fovea broad, clearly occupying more than half of the distance
	between the lateral ocellus and the compound eye (Fig. 37C). Terga with
	strong apical pale hair bands that uniformly cover the entirety of the tergal
	depressions and obscure the underlying surface (Figs 37D, 38D; North Af-
	rica to Israel)
_	Fovea narrow, occupying at most half the distance between the lateral ocel-
	lus and the compound eye (Figs 33B, 36C). Terga with or without clear hair

	bands; if with hair bands, then hair bands typically do not uniformly cover
	the tergal depressions, either weaker basally or stronger apically3
3	T1 with extremely sparse punctures, punctures separated by 2–4 puncture
	diameters. Facial fovea along its entire length separated from the inner mar-
	gin of the compound eye by a distance equal to its own diameter (south-
	western Morocco only) sparsipunctata Wood
_	T1 with dense punctures, punctures separated at most by 2 puncture diam-
	eters, usually by 1 puncture diameter (Fig. 59D). Fovea not strongly sepa-
	rated from the inner margin of the compound eye4
4	Pubescence of scutum weakly squamous, anterior dorsolateral corners of
•	scutum with pubescence longer, clearly exceeding width of antennae in
	length. Terga always predominantly red. Terga with clear apical hair bands.
	Punctation of T1 slightly spaced, punctures separated by 1–2 puncture di-
	ameters (eastern and south-eastern Spain only)
_	Pubescence of scutum strongly squamous and short, anterior dorsolat-
	eral corners of scutum with pubescence short, clearly shorter than width
	of antennae (Figs 33C, 59C). Terga variable, red to black or any inter-
	mediate combination. Terga with or without clear apical hair bands.
	Punctation of T1 denser, punctures separated at most by 1 puncture
	diameter5
5	A3 exceeding A4+5 in length. Terga always extensively red-marked (central
	and south-eastern Spain)parata Warncke
_	A3 equalling A4+5 in length. Terga variable, from almost entirely black to
	entirely red-marked6
6	Terga usually predominantly dark (Figs 33D, 36D). Found in Spain
_	Terga always extensively or entirely red-marked (Figs 35D, 59D). Found
	outside of Spain
7	Clypeus at least partly yellow-marked (North Africa to the Middle East)
	succinea Dours
_	Clypeus uniformly dark8
8	A3 very short, shorter than A4 (North Africa to Israel) caroli Pérez
_	A3 at least slightly longer than A49
9	A3 exceeding A4+5 in length (central and south-eastern Spain)
_	A3 not exceeding A4+510
10	Genital capsule without clear kink in the inner margins of the gonostyli
10	(Fig. 85E). Scutum less densely punctate, punctures separated by 0.5 punc-
	ture diameters, with shiny interspaces, predominantly shiny (eastern and
	south-eastern Spain only)
	Genital capsule with clear kink in the inner margins of the gonostyli
_	
	(Fig. 85F). Scutum more densely punctate, punctures almost confluent,
	predominantly dull (central and southern Spain) pruinosa Erichson

Description of new species

Andrena (Avandrena) juliae Wood, sp. nov.

https://zoobank.org/C53C51E7-AA4A-43A1-85CC-6C4862ED1D19

Type material. *Holotype***. SPAIN:** Cádiz, Parque Natural Los Alcornocales, Las Algamitas, Finca Murtas, 36.3273°N, -5.5986°W, 18.iii.2023, 1, leg. T.J. Wood, OÖLM.

Paratypes. Spain: Cádiz, Parque Natural Los Alcornocales, Las Algamitas, Finca Murtas, 18.iii.2023, 1♀, leg. T.J. Wood, TJWC; Cádiz, Tarifa, 1 km N, grazing fields, 23.iii.2023, 1♀, leg. T.J. Wood, OÖLM.

Description. Female. Body length: 9 mm (Fig. 47A). *Head:* Dark, 1.4 times wider than long (Fig. 47B). Clypeus weakly domed, with fine granular shagreen, dull, overlain by weak raised rugosity, rugosity forming weak, obscure, and interrupted raised latitudinal striations; remaining surface impunctate. Process of labrum rounded rectangular, twice as broad as long, surface smooth and shiny. Gena equalling width of compound eye; ocelloccipital distance equals diameter of lateral ocellus. Foveae moderately broad, occupying half space between compound eye and lateral ocellus, short, ventrally extending to level of antennal insertions; foveae filled with black hairs (Fig. 47C). Face medially with light brown-white hairs covering clypeus and antennal insertions, intermixing with black hairs, becoming entirely black haired along inner margin of compound eyes and frons; gena and vertex with long light brown hairs, longest hairs equalling length of scape. Antennae dark, A7–12 ventrally slightly lightened grey by presence of scales; A3 equalling A4+5+6. *Mesosoma*: Scutum and scutellum with fine granular shagreen, very weakly shiny, predominantly dull, underlying surface shallowly and obscurely punctate, punctures separated by 1–2 puncture diameters (Fig. 47D). Pronotum rounded. Mesepisternum and dorsolateral parts of propodeum with fine granular shagreen, predominantly dull, shagreenation overlain by fine network of raised rugosity, not forming a linked network. Propodeal triangle short and broad, slightly depressed below level of dorsolateral parts of propodeum, with extremely fine granular shagreen, basal 2/3rds covered with raised longitudinal rugae, propodeal triangle thus strongly contrasting dorsolateral parts of propodeum (Fig. 47E). Mesepisternum with long strongly plumose hairs, hairs predominantly pale with some intermixed black hairs, hairs clearly exceeding length of scape. Scutum and scutellum with strongly plumose light brown hairs, intermixed with black hairs medially. Propodeum with long strongly plumose light brown hairs, propodeal corbicula incomplete, dorsal fringe weak and poorly defined, internal surface with numerous pale simple hairs. Legs dark, tarsi obscurely lightened reddish brown; pubescence light brown. Flocculus complete, strong, composed of strongly plumose light brown hairs; femoral and tibial scopa composed of light brown simple hairs. Hind tibial claws with strong inner tooth. Wings hyaline, stigma and venation dark orange, nervulus interstitial. *Metasoma*: Terga dark, apical rim of marginal areas narrowly lightened hyaline-yellow (Fig. 47F). Tergal discs with scattered and small hair-bearing punctures, punctures separated by 3-4 puncture diameters, underlying surface finely shagreened, silky smooth, shiny. Tergal discs with

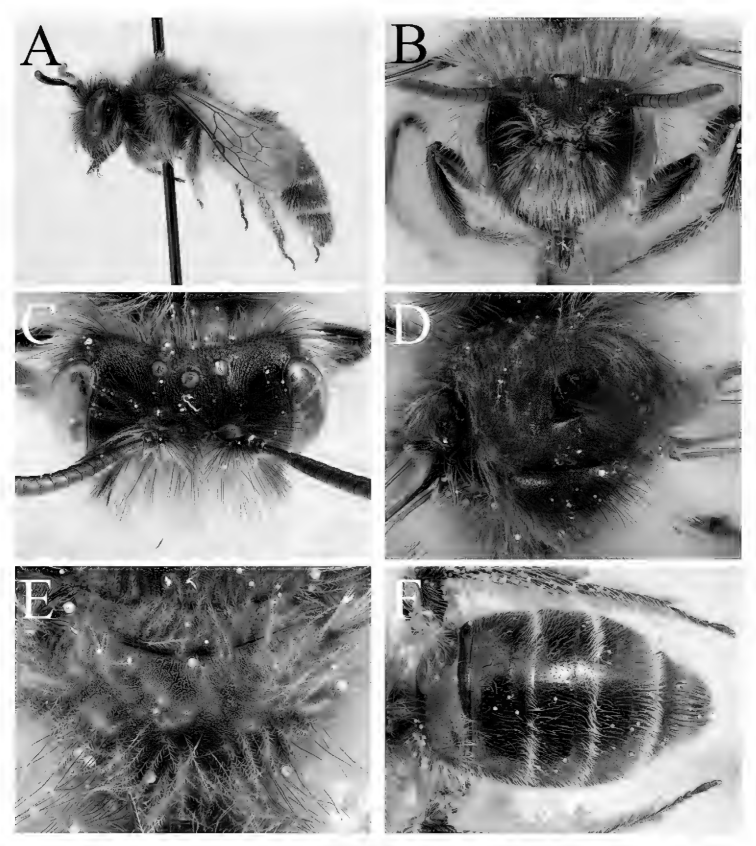


Figure 47. Andrena (Avandrena) juliae sp. nov. female **A** profile **B** face, frontal view **C** head, dorsal view **D** scutum, dorsal view **E** propodeum, dorsal view **F** terga, dorsal view.

sparse upstanding pale hair, T2–4 with dense apical whitish hair bands that obscure underlying surface. Apical fringe of T5 and hairs flanking pygidial plate dark brown; pygidial plate apically rounded, surface flat and featureless.

Male. Unknown.

Diagnosis. Andrena juliae can be recognised as belonging to the subgenus Avandrena due to its moderate to small body size (9 mm), short and wide head (clearly wider than long), and short and wide foveae that are only slightly longer than wide, as well as the behavioural observation that it is strongly associated with *Erodium* (Geranicaceae,

see Remarks). The posterior face of the hind femora lacks spines, separating the species from *A. avara* Warncke, 1967 and *A. panurgina* De Steffani, 1889 and placing it close to *A. melacana* Warncke, 1967 and *A. erodiorum* Wood & Ortiz-Sánchez, 2022.

Andrena juliae can be separated from both species by the structure of the propodeum which has the dorsolateral parts with fine granular shagreenation that is overlain by a fine network of raised rugosity, this rugosity not forming a linked network. The propodeal triangle itself is clearly differentiated, slightly depressed below the level of the surrounding parts of the propodeum and delineated by fine carinae, the surface with fine granular shagreenation and with a network of raised longitudinal carinae covering the basal 2/3rds. In contrast, A. erodiorum has the dorsolateral parts of the propodeum shiny, overlain with a fine network of raised rugosity that joins together to form a clear network (Fig. 73E). The propodeal triangle is clearly delinated by raised carinae and with the entire surface covered with longitudinal carinae. In A. melacana, the dorsolateral parts of the propodeum are entirely covered with fine granular shagreen, without an overlay of raised rugosity, with the propodeal triangle not depressed and with even finer granular shagreen, not strongly differentiated from the surrounding areas.

The propodeum of *A. juliae* therefore sits between both comparison species and is distinct from both. It can be further separated by the pubescence of the mesepisternum which is predominantly composed of pale hairs, with approximately 30% of these hairs black (in *A. melacana* with 50–60% of the hairs of the mesepisternum black; in *A. erodiorum* with only 10% of these hairs black), by the facial foveae which occupy half of the space between the compound eye and a lateral ocellus (occupying ¾ of this space in *A. erodiorum*), by the colour of the hairs of the apical fringe of T5 and those flanking the pygidial plate which are dark brown (golden-brown in *A. erodiorum*, dark brown in *A. melacana*), and by the pubescence of the terga which are covered in sparse erect white hairs, T2–4 with dense apical hair bands of white hairs that obscure the underlying surface (in *A. melacana* terga with sparse short pubescence, only forming weak apical tergal hair bands that do not obscure the underlying surface; pubescence very similar in *A. erodiorum*).

Remarks. All specimens were collected from *Erodium* spp. The two females from Las Algamitas were collecting pollen from this genus (*Erodium* pollen can be seen in Fig. 47A). Across sampling locations in Cádiz province during March 2023, four species of *Avandrena* were active, specifically *A. avara* sensu stricto, *A. juliae*, *A. melacana*, and *A. panurgina*. All species appear to be narrowly oligolectic on *Erodium* (see below), and *A. juliae* could be found in direct sympatry with *A. avara* sensu stricto, *A. melacana*, and *A. panurgina* at the Las Algamitas site and with *A. melacana* and *A. panurgina* at the Tarifa site. *Andrena erodiorum* is currently known only from south-eastern Spain (Albacete). One *A. juliae* female showed abraded body hairs, and no males were captured, implying that the flight season may begin in early March or even late February, perhaps offering a reason as to why this species has been overlooked until now.

Etymology. Dedicated to my friend and colleague Julia Jones (University College Dublin, Ireland) who invited me on the University field course during which this new species was discovered.

Distribution. Spain (Cádiz province).

Other material examined. *Andrena avara* s. str.: Spain: Cádiz, Parque Natural Los Alcornocales, Las Algamitas, Finca Murtas, 18.iii.2023, 1, leg. T.J. Wood, TJWC; Cádiz, Bolonia, El Lentiscal, 24.iii.2023, 1, leg. T.J. Wood, TJWC.

Andrena melacana: Spain: Cádiz, Parque Natural Los Alcornocales, Las Algamitas, Finca Murtas, 18.iii.2023, 3&, 2\$\, leg. T.J. Wood, TJWC; Cádiz, Tarifa, 1 km N, grazing fields, 19.iii.2023, 4&, 6\$\, leg. T.J. Wood, TJWC.

Andrena panurgina: Spain: Cádiz, Parque Natural Los Alcornocales, Las Algamitas, Finca Murtas, 18.iii.2023, 4♂, 3♀, leg. T.J. Wood, TJWC; Cádiz, Tarifa, 1 km N, grazing fields, 19–23.iii.2023, 6♂, 2♀, leg. T.J. Wood, TJWC.

Andrena (Euandrena) isolata Wood, sp. nov.

https://zoobank.org/08390D0E-AC36-4859-AE7E-BDD126D34BB8

Type material. *Holotype.* **SPAIN:** Granada, Sierra Nevada, Trevélez to Refugio La Campiñuela, 37.0239°N, -3.2656°W, 1700–2400 m, 14.vi.2021, 1\$\bigcap\$, leg. T.J. Wood, OÖLM [BOLD accession number WPATW368-21].

Description. Female. Body length: 9.5 mm (Fig. 48A). *Head*: Dark, 1.2 times wider than long (Fig. 48B). Clypeus weakly domed, densely and evenly punctate,

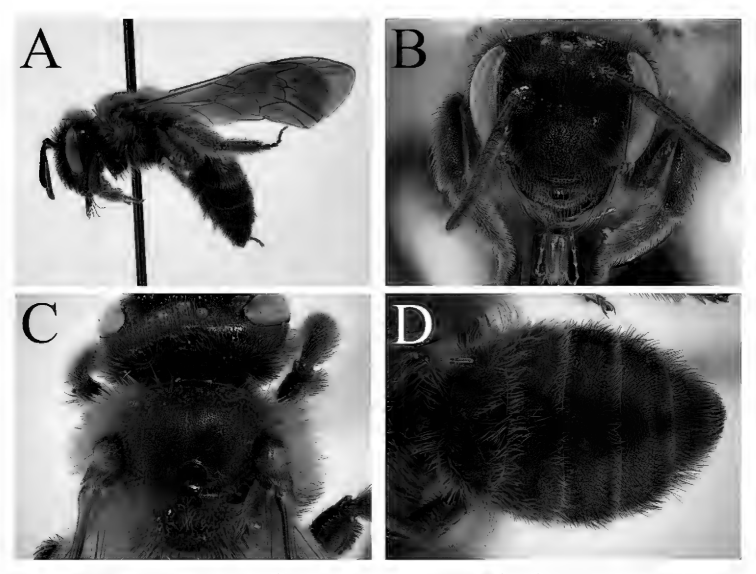


Figure 48. Andrena (Euandrena) isolata sp. nov. female **A** profile **B** face, frontal view **C** scutum, dorsal view **D** terga, dorsal view.

punctures separated by <0.5 puncture diameters, interspaces shiny. Process of labrum trapezoidal, twice as broad as long, apical margin with very weak emargination medially. Gena slightly exceeding width of compound eye; ocelloccipital distance 1.5 times diameter of lateral ocellus; vertex behind ocellar triangle densely punctate, punctures separated by 0.5 puncture diameters. Fovea dorsally narrow, occupying 1/3 of space between lateral ocellus and compound eye, ventrally reaching level of antennal insertions, here narrowing to ½ dorsal width; fovea filled with black hairs. Face, gena, and scape covered with uniformly short black hairs, vertex dorsally with predominantly golden-brown hairs with scattered intermixed black hairs. Antennae dark, A3 slightly exceeding A4+5, shorter than A4+5+6. *Mesosoma*: Scutum and scutellum densely punctate, punctures predominantly separated by <0.5 puncture diameters, at most by 1 puncture diameter medially, underlying surface finely shagreened and weakly shiny (Fig. 48C). Pronotum rounded. Mesepisternum microreticulate, weakly shiny to dull, with obscure raised reticulation. Dorsolateral parts of propodeum microreticulate, weakly shiny, densely and shallowly punctate, punctures separated by 0.5–1 puncture diameter. Propodeal triangle narrow, internal surface finely granulate and impunctate, basally with obscurely and weakly raised rugosity, propodeal triangle thus strongly contrasting punctate dorsolateral parts of propodeum. Mesepisternum ventrally with moderately long black hairs, becoming intermixed with golden-brown hairs medially, with entirely golden-brown hairs dorsally, hairs at most equalling length of scape. Scutum and scutellum dorsally with short golden-brown hairs, scutum medially with very short and obscure black hairs. Propodeum with long golden-brown hairs, propodeal corbicula incomplete, dorsal fringe composed of long plumose golden-brown hairs, internal surface with numerous plumose light hairs, becoming darker ventrally. Legs predominantly dark, hind femorae ventrally and posterior face of hind tibiae obscurely lightened reddish-orange; pubescence black to orange-brown. Flocculus incomplete, short, composed of weakly plumose orange hairs; femoral and tibial scopa orange, tibial scopa with at most occasional brown hairs dorso-basally. Hind tibial claws with strong inner tooth. Wings weakly infuscate, stigma and venation dark brown, nervulus interstitial. Metasoma: Terga dark, apical rim of marginal areas very narrowly lightened hyaline-brown (Fig. 48D). Tergal discs regularly punctate, punctures separated by 1-2 puncture diameters, underlying surface shagreened, weakly shiny; punctures extending onto marginal areas, here weak and obscure, separated by 2-3 puncture diameters. T1-2 with orange-brown hairs basally, laterally, and apically; T2-3 with orange-brown hairs forming weak apical hair bands, not obscuring underlying surface, T4 with weak apical hair band black; discs of T3-4 with short black hairs. Apical fringe of T5 and hairs flanking pygidial plate black; pygidial plate rounded triangular, with slightly depressed and densely punctate area medially, lateral margins impunctate.

Male. Unknown.

Diagnosis. Andrena isolata can be quickly recognised as a Euandrena due to the narrow facial foveae (dorsally occupying ½ of space between the lateral ocellus and the compound eye) which narrow further ventrally combined with the long A3

(slightly exceeding length of A4+5) and the simple, non-plumose hairs of the tibial scopae. Its generally dark appearance with orange-brown hairs dorsally on the mesosoma and tibial scopa plus at least some black hairs on the mesepisternum place it immediately close to *A. bicolor* and allied taxa. As discussed above, the subgenus *Euandrena* is taxonomically complex, and multiple genetically distinct taxa have been lumped under *A. bicolor*. In an Iberian context, *A. isolata* is best diagnosed against *A. bicolor* s.l., *A. fortipunctata* Wood, 2021, and the distinct and probably undescribed taxon in north-western Africa identified above. Extreme care should be taken when identifying specimens morphologically, and barcodes should be used whenever possible.

In direct comparison to barcoded A. bicolor s.l. specimens, the only consistent character that can be identified is the structure of the clypeus. Andrena isolata has the clypeus densely punctate, with punctures separated by <0.5 puncture diameters, interspaces shiny but overall the clypeus only weakly shiny due to the small size of the interspaces (Fig. 48B). In A. bicolor s.l., the clypeal punctures are separated by an average of at least 1 puncture diameter, sometimes more, and the clypeus is therefore more strongly shiny due to the larger interspaces. The dense clypeal punctures place A. isolata close to A. fortipunctata, but the two species can be separated by the tergal structure (see illustrations in Wood et al. 2021), with the tergal margins weakly depressed with at most the apical rim lightened hyaline yellow (tergal margins strongly depressed and extensively lightened semi-translucent brown in A. fortipunctata), terga punctate with some punctures extending onto tergal margins, here separated by 2-3 puncture diameters (tergal margins impunctate in A. fortipunctata), margins of T2-3 with weak light brown hair fringes apically, T4 with entirely black hairs (T2-4 with white hair fringes emerging from junction between disc and margin, these hair fringes overlying marginal areas in A. fortipunctata), terminal fringe of T5 and hairs flanking pygidial plate black (terminal fringe and hairs flanking pygidial plate dark brown in A. fortipunctata). Andrena isolata is almost identical to the probably undescribed taxon from northwestern Africa, but can also be separated by the structure of the clypeus. Andrena isolata has the clypeus densely punctate with punctures separated by <0.5 puncture diameters and with the interspaces shiny, whereas in the taxon from north-western Africa, the punctures are equally dense but the interspaces are shagreened and dull. Furthermore, this latter taxon has a short and shallow longitudinal furrow at the apex of the clypeus that is absent in A. isolata.

Remarks. Andrena isolata probably represents a relictual species that has become isolated on the Sierra Nevada from what is now a remaining North African population. Additional genetic sampling is needed to establish whether it is found away from the Sierra Nevada, but a specimen from the nearby Sierra de Baza collected at an altitude of 2000 m barcoded as A. bicolor s.l. [WPATW297-21]. Andrena isolata may well be restricted to the Sierra Nevada. Additional sampling is required to establish its ecology, including its voltinism. Its capture on Campanula implies that it has a similar ecology to A. bicolor s.l. (see Praz et al. 2019), but this requires dedicated study.

Etymology. Derived from the Latin *insulatus*, to be made into an island, *isolata* (feminine form) thus means to be isolated, in reference to its presence on the Sierra Nevada, separated from its nearest genetic relative in North Africa.

Distribution. Spain (Sierra Nevada).

Andrena (Micrandrena) ortizi Wood, sp. nov.

https://zoobank.org/F6956353-0278-4AE3-8C48-4084EDC29CCC

Type material. *Holotype.* **S**PAIN: Granada, Sierra Nevada, Mirador Monte Ahí de Cara, 37.1239°N, -3.4322°W, 2100 m, 6.vi.2021, 1\(\sigma\), leg. T.J. Wood, on *Vella spinosa* (Brassicaceae), OÖLM [BOLD accession number WPATW972-22].

Paratypes. Spain: Granada, Sierra Nevada, Mirador Monte Ahí de Cara, 2100 m, 6.vi.2021, 2♀, leg. T.J. Wood, on *Vella spinosa* (Brassicaceae), OÖLM/TJWC; Granada, Sierra Nevada, Puerto de la Ragua, Barranco Maja Caco, 2000 m, 10.vi.2021, 1♀, leg. T.J. Wood, TJWC; Granada, Sierra Nevada 1900 m, ri. Veleta, 1.vi.1982, 1♀, leg. R. Leys, RMNH; Granada: Órgiva N, 1300 m, 26.vi.1988, 1♀, leg. M. Schwarz, OÖLM.

Description. Female. Body length: 7 mm (Fig. 49A). *Head*: Dark, 1.1 times wider than long (Fig. 49B). Clypeus long, apical margin clearly ventrally exceeding level of line between lower margins of compound eyes, surface weakly domed, irregularly punctate, punctures separated by 1-3 puncture diameters, impunctate longitudinal midline present; underlying surface basally finely shagreened and weakly shiny, shagreenation becoming weaker apically, here almost smooth and shiny. Process of labrum narrowly trapezoidal, slightly broader than long, apical margin narrowly truncate. Gena slightly exceeding width of compound eye; ocelloccipital distance equalling diameter of lateral ocellus. Fovea dorsally narrow, occupying 1/4 space between lateral ocellus and compound eye, ventrally extending below level of antennal insertions, consistently wide along its length, filled with white hairs. Face, gena, vertex, and scape covered with sparse whitish hairs, none equalling length of scape. Antennae dark, A3 equalling length of A4+5. *Mesosoma*: Scutum and scutellum regularly punctate, punctures separated by 1 puncture diameter, underlying surface shagreened and weakly shiny (Fig. 49C). Pronotum rounded. Mesepisternum microreticulate, weakly shiny with finely raised reticulation. Dorsolateral parts of propodeum with dense network of reticulation; propodeal triangle laterally delineated with raised carinae, internal surface with dense network of raised rugosity. Mesepisternum, scutum, and scutellum with long whitish hairs, none equalling length of scape. Propodeal corbicula incomplete, dorsal fringe composed of long whitish plumose hairs, internal surface with scattered long simple whitish hairs. Legs dark, pubescence whitish to light brownish. Flocculus complete, composed of short weakly upturned plumose hairs; flocculus, femoral and tibial scopae white, tibial scopae with some brown hairs dorso-basally. Hind tibial claws with inner tooth. Wings hyaline, stigma dark brown, venation brown, nervulus interstitial. Metasoma: Terga dark, apical rim of marginal areas very narrowly lightened hyalinebrown. Tergal discs densely and clearly punctate, disc of T1 with punctures separated



Figure 49. *Andrena (Micrandrena) ortizi* sp. nov. female **A** profile **B** face, frontal view **C** scutum, dorsal view **D** terga, dorsal view.

by 1 puncture diameter, T2–3 with punctures separated by 0.5 puncture diameters, underlying surface shagreened and weakly shiny (Fig. 49D). Marginal areas impunctate, with rectangular shagreen; marginal areas long, on T1 occupying ¼ of tergum, on T2 occupying ½ of tergum, on T3 occupying 2/3rds of tergum, on T4 occupying almost all visible tergum. T2–4 with long sparse plumose hairs arising from base of marginal area, covering but not obscuring marginal area; T3–4 apically with dense short apical fringe of white hairs laterally, obscuring underlying surface. Apical fringe of T5 and hairs flanking pygidial plate dark brown with occasional whitish hairs laterally; pygidial plate rounded triangular, with obscurely raised medial ridge, otherwise featureless.

Male. Unknown.

Diagnosis. Andrena ortizi can quickly be recognised as a Micrandrena due to its small body size, dark integument, and entirely rugose propodeal triangle. Due to the comparatively (for a Micrandrena) long face and clypeus (head overall only 1.1 times wider than broad; apical margin of clypeus clearly ventrally exceeding level of a line drawn between the lower margins of the compound eyes), narrow facial fovea (dorsally occupying ¼ of space between the lateral ocellus and the compound eye, consistently wide along its length, and densely punctate scutum (punctures separated by <1 puncture diameter) they are comparable to A. rugulosa Stöckhert, 1935 (Switzerland to Lebanon

and the Caucasus), *A. atlantea* Wood, 2021 (High and Middle Atlas Mountains in Morocco), and an undescribed *Micrandrena* species from the Middle Atlas (see above).

Andrena ortizi can be separated from A. rugulosa by the scutum which is slightly less densely punctate, punctures separated by 1 puncture diameter (punctures separated by 0.5 puncture diameters in A. rugulosa), the underlying surface of the scutum being finely shagreened and shiny (scutum is densely shagreened and dull in A. rugulosa), the tergal discs are strongly and clearly punctate (tergal discs obscurely punctate in A. rugulosa), and the marginal areas of T2-4 occupy at least ½ the visible length of the tergum, on T3-4 clearly occupying over ½ this length (marginal areas typically occupying ½ length of tergum, at most occupying ½ tergum on T4 in A. rugulosa). Andrena ortizi is more similar to A. atlantea, sharing a similarly less densely punctate and weakly shiny scutum, but the same character of wide tergal margins can be used to separate them, with the marginal areas occupying at most $\frac{1}{3}$ of the length of the tergum in A. atlantea. Finally, A. ortizi is most similar to the undescribed Micrandrena from the Middle Atlas, and the width of the tergal margins can again be used to separate them, with the tergal margins occupying at most 1/3 of the disc of T2 and 1/2 of the discs of T3-4. Additionally, A. ortizi has the disc of T2-3 clearly and densely punctate, whereas in the undescribed Micrandrena the discs of T2-3 are at most obscurely punctate, with punctures disappearing into the background microreticulation. Andrena ortizi also has a strongly isolated distribution, separated from the Swiss Alps (A. rugulosa) by c. 1,300 km and the high altitude parts around Ifrane and Azrou in the Middle Atlas (A. atlantea, the undescribed Micrandrena species) by c. 400 km.

Remarks. At the Mirador Monte Ahí de Cara (Fig. 50A, see also Fig. 22A), this species was collected foraging for pollen on the spiny yellow species *Vella spinosa* (Brassicaceae) in open habitat just above the tree line. *Andrena ortizi* appears to be endemic to the Sierra Nevada. Searches at elevations lower than 2000 m did not detect the species during 2021 surveys, so the collection nominally made at 1300 m near Órgiva may not be representative – the slopes above Órgiva (southern slopes of the Sierra Nevada) ascend rapidly, the collector may well have climbed to a higher altitude when collecting this specimen. The morphological similarity between *A. rugulosa*, *A. ortizi*, and *A. atlantea* suggests a pattern of geographical isolation in montane habitats, though only *A. ortizi* and *A. atlantea* show a close genetic relationship based on the COI sequences.

Etymology. Dedicated to the Spanish naturalist and hymenopterist Francisco Javier Ortiz-Sánchez who has worked extensively on the Iberian bee fauna for many years, including that of the Sierra Nevada.

Distribution. Spain (Sierra Nevada).

Andrena (Truncandrena) ghisbaini Wood, sp. nov.

https://zoobank.org/E5ABB0AB-EE49-4CCC-91B4-9A9A99AD2A5C

Type material. *Holotype.* **S**PAIN: Málaga, PN Sierra de las Nieves, mountain peak S of Pinsapo Escalereta, 36.6621°N, -5.0362°W, 1600 m, 30.v.2021, 1\(\text{Q}\), leg. T.J. Wood, OÖLM [BOLD accession number WPATW239-21].

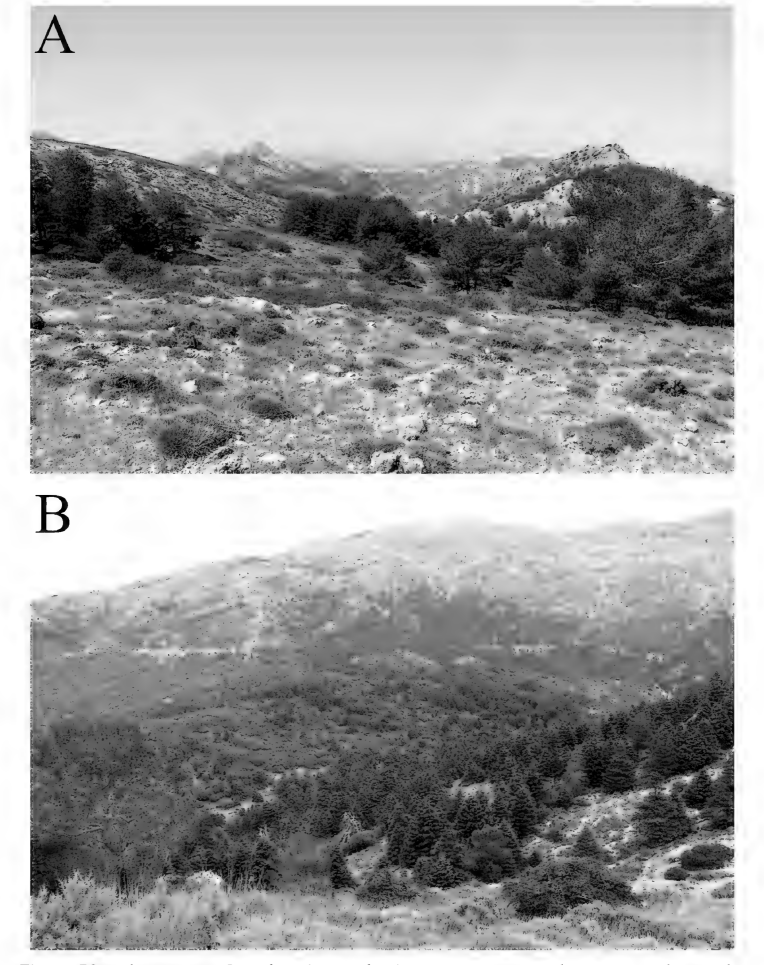


Figure 50. Habitat context **A** *Andrena* (*Micrandrena*) *ortizi* sp. nov., Granada, Sierra Nevada, Mirador Monte Ahí de Cara, 2100 m, 12.vi.2021 **B** *Andrena* (*Truncandrena*) *ghisbaini* sp. nov., Málaga, PN Sierra de las Nieves, mountain peak S of Pinsapo Escalereta, 1600 m, 30.v.2021.

Paratypes. Spain: Málaga, PN Sierra de las Nieves, mountain peak S of Pinsapo Escalereta, 1600 m, 30.v.2021, 1♀, leg. G. Ghisbain, TJWC; Málaga – Elvira, 11.ii.1981, 4♂, leg. H. Teunissen, RMNH.

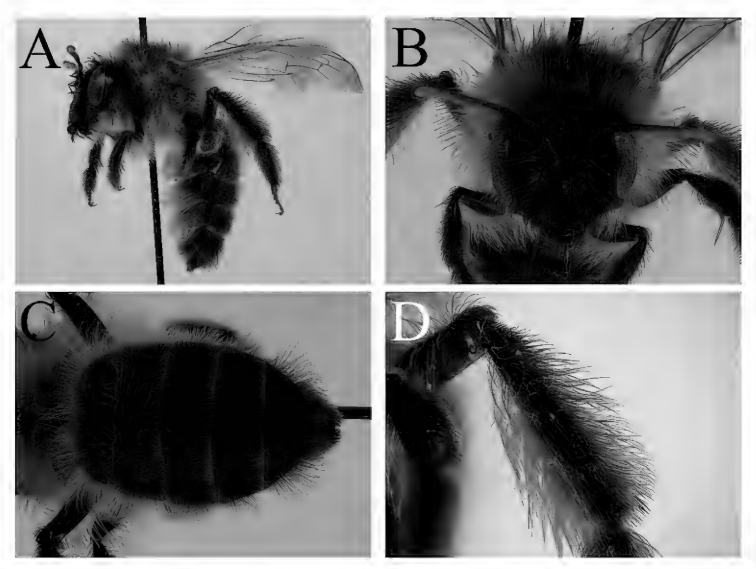


Figure 51. Andrena (Truncandrena) ghisbaini sp. nov. female **A** profile **B** face, frontal view **C** terga, dorsal view **D** tibial scopa, profile view.

Description. Female. Body length: 15–16 mm (Fig. 51A). *Head*: Dark, 1.2 times wider than long (Fig. 51B). Clypeus weakly domed, clearly punctate, punctures separated by 0.5–1 puncture diameters with exception of median longitudinal impunctate line, narrow basally, broadening apically, thus elongate triangular; underlying surface finely shagreened, weakly shiny. Process of labrum broadly trapezoidal, twice as broad as long, apical margin clearly emarginate. Gena broad, almost two times width of compound eye; ocelloccipital distance 1.5 times diameter of lateral ocellus. Fovea dorsally occupying slightly more than ½ space between lateral ocellus and compound eye, extending ventrally to lower margins of antennal insertions, filled with black hairs. Gena ventrally and laterally with long light brown hairs, longest equalling length of scape, hairs becoming black on vertex, scape, and majority of face, with shorter light brown hairs around antennae insertions. Antennae basally dark, A4 apically, A5–12 ventrally extensively lightened orange; A3 exceeding A4+5, shorter than A4+5+6. Mesosoma: Scutum and scutellum with extremely shallow and obscure punctures, punctures separated by 0.5-1 puncture diameters, punctures disappearing into underlying fine granular shagreen, surface dull to very weakly shiny. Pronotum rounded. Mesepisternum and dorsolateral parts of propodeum with fine granular shagreen, weakly shiny, with fine and scattered raised hair-bearing punctures, punctures separated by 2-3

puncture diameters; propodeal triangle broad, with extremely fine granular shagreen, without hair-bearing punctures, propodeal triangle thus defined by change in surface sculpture from dorsolateral parts of propodeum. Mesepisternum and propodeum with long finely plumose light brown hairs, clearly exceeding length of scape; scutum and scutellum medially with long black hairs and occasional isolated pale hairs covering majority of disc, laterally becoming intermixed with light brown hairs. Propodeal corbicula incomplete, very weakly defined, dorsal fringe not differentiated from hairs of internal surface, both parts composed of long finely plumose long light brown hairs. Legs dark, apical tarsal segments lightened dark reddish, pubescence dark brown. Flocculus complete, composed of long weakly plumose and upturned light brown hairs; femoral scopae composed of light brown simple hairs; tibial scopa long, hairs exceeding apical width of hind tibia, hairs dorsally dark brown, ventrally golden orange (Fig. 51D). Hind tarsal claws with inner tooth. Wings hyaline, stigma dark brown, venation dark brown to orange, nervulus interstitial. Metasoma: Terga dark, apical rim of marginal areas narrowly lightened hyaline-yellow; discs with extremely fine granular shagreen, weakly shiny, with fine and scattered hair-bearing punctures, punctures separated by 3–4 puncture diameters (Fig. 51C). Disc of T1 with long light brown hairs, exceeding length of scape, decreasing in length over T2–3, disc of T3 with intermixed short light brown and black hairs, becoming predominantly black on discs of T4-5. T2-4 with weak apical fringes of short light brown hairs, not obscuring underlying surface. Apical fringe of T5 and hairs flanking pygidial plate dark brown, pygidial plate rounded triangular, with weakly raised medial area, otherwise featureless.

Male. Body length: 13–14 mm (Fig. 52A). *Head*: Dark, 1.3 times wider than long (Fig. 52B). Clypeus weakly domed, entirely yellow-marked with exception of two dark rounded spots medio-laterally. Clypeus punctured, punctures separated by 0.5-1 puncture diameters with exception of median longitudinal impunctate line, essentially non-existent basally, broadening apically, thus elongate triangular; underlying surface finely shagreened, weakly shiny. Process of labrum broadly rectangular, 2.5 times wider than long, apical margin weakly emarginate, surface smooth and shiny. Gena broad, 2 times width of compound eye; ocelloccipital distance 1.5 times diameter of lateral ocellus. Gena ventrally and laterally with long light brown hairs clearly exceeding length of scape, becoming intermixed with black hairs on vertex. Face medially with extensive whitish to light-brownish hairs on clypeus, antennae insertions, and scape intermixed with black hairs along inner margins of compound eyes and scape. Antennae basally dark, A4–13 ventrally lightened orange; A3 longer than A4, shorter than A4+5; A4 rectangular, longer than broad, slightly shorter than A5. *Mesosoma*: Mesosoma structurally as in female; pubescence as in female. Legs basally dark, apical tarsal segments and hind tibiae lightened dark reddish-brown, pubescence dark brown to orange brown. Hind tarsal claws with inner tooth. Wings hyaline, stigma orange, venation dark brown to orange, nervulus interstitial. *Metasoma*: Terga structurally as in female. Discs of T1-4 with long light brown hair, on T1 clearly exceeding length of scape, becoming progressively shorter to T4; T5-6 with short black hairs on disc (Fig. 52C). T2-4 with weak apical hair fringes apically, not

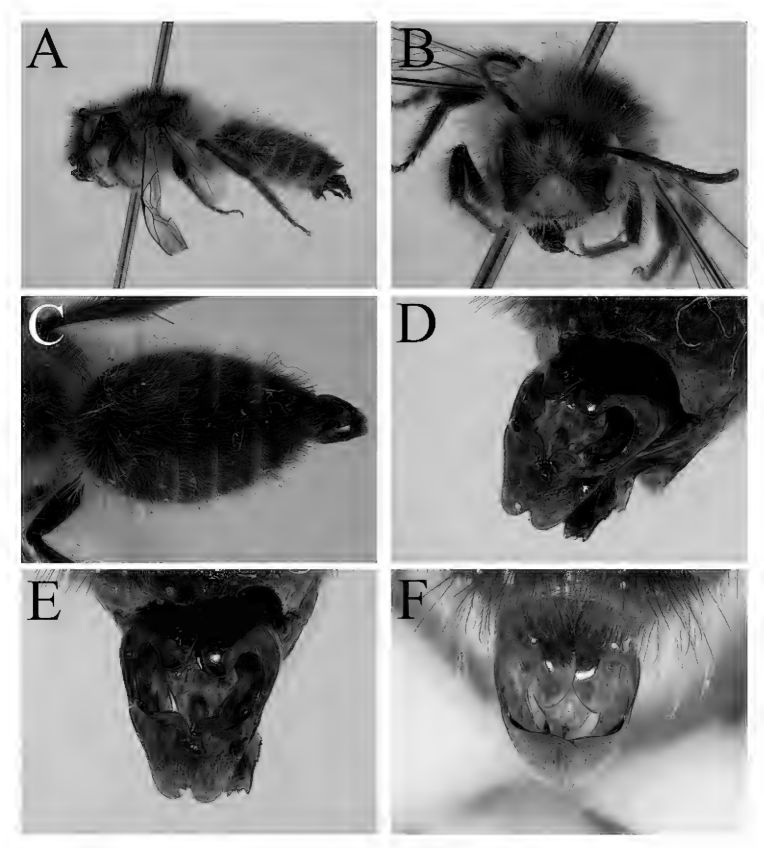


Figure 52. Andrena (Truncandrena) ghisbaini sp. nov. male **A** profile **B** face, frontal view **C** terga, dorsal view **D** genital capsule, dorsolateral view **E** genital capsule, dorsal view; Andrena (Truncandrena) villipes Pérez, 1895 male **F** genital capsule, dorsal view.

obscuring underlying surface. S8 columnar, apex rounded, ventral surface covered with short brown hairs. Genital capsule slightly elongate, gonocoxae produced into strong apical teeth, teeth pointed with apical margins diverging (Fig. 52D, E). Gonostyli basally narrow, apically produced and flattened into rounded triangular plates, internal margin strongly raised and reflexed. Penis valves basally broad, occupying more than ½ space between gonostyli, with narrow laterally produced hyaline extensions; penis valves strongly narrowing apically.

Diagnosis. Andrena ghisbaini can be recognised within Truncandrena due to its characteristically smooth and finely granulate propodeal triangle which contrasts with the similarly granulate dorsolateral parts of the propodeum which bear fine and scattered raised hair-bearing punctures, the rounded pronotum, the linear malar space, the large body size (>13 mm), yellow male clypeus, and typical genital capsule with the inner margins of the flattened apical parts of the gonostyli strongly raised. It can be placed closest to A. villipes Pérez, 1895 (Fig. 30) due to the antennae that are ventrally extensively lightened orange, the intermixed light and dark pubescence of the face, with pale hairs medially and dark hairs laterally, the intermixed light and dark pubescence of the scutum, with dark hairs medially and lighter hairs laterally, by the entirely yellow-marked male clypeus, and by the penis valves basally broad with lateral hyaline extensions, valves strongly narrowing apically.

The immediate difference between the two taxa is size, with *A. villipes* averaging 12–13 mm in length in females and 11–12 mm in males, compared to 15–16 mm and 13–14 mm respectively in *A. ghisbaini*. Structurally, *A. ghisbaini* females can be separated by the bicoloured scopa, black dorsally and orange ventrally (Fig. 51D; *A. villipes* with scopa unicolourous orange, Fig. 30B), the longer face, clypeus clearly projecting ventrally well below a line drawn between the lower margins of the compound eyes (*A. villipes* with face shorter, clypeus only slightly projecting below this line in direct comparison), the clypeus clearly punctured with a longitudinal impunctate midline that broadens apically (*A. villipes* with clypeus obscurely and shallowly punctate, without obvious impunctate midline), and the reduced pale pubescence of the face, with light hairs restricted to the area around the antennal insertions (*A. villipes* with extensive pale hairs covering majority of face and clypeus, with black hairs predominantly along the inner margins of the compound eyes).

In the male sex, *A. ghisbaini* can be separated by the same clypeal punctation character (stronger in *A. ghisbaini* with clearer impunctate midline), but this is slightly more subtle than in the female sex. Direct comparison of the genital capsule shows that flattened apical part of the gonostyli are more strongly elongate and longer than broad, thus appearing triangular (Fig. 52D, E; in *A. villipes* with the flattened apical part of the gonostyli more rounded, about as long as broad, Fig. 52F), the inner margins of these parts more strongly and acutely raised, slightly reflexed (in *A. villipes* with the inner margin less strongly raised and not reflexed).

Remarks. The two females from the Sierra de las Nieves (Fig. 50B) were collected from *Cistus albidus* Linnaeus (Cistaceae). Like *A. villipes* (Table 1), this species is likely to be oligolectic on Cistaceae. Additional surveys are needed to clarify the limits of its range. Male specimens from Elvira in the Naturalis collection were incorrectly determined by Teunissen as *A. maroccana* Benoist, 1950 which is a synonym of *A. leptopyga* Pérez, 1895.

Etymology. Dedicated to my friend and colleague Guillaume Ghisbain (Mons, Belgium) who accompanied me during fieldwork in Málaga province, and who is an accomplished hymenopterist in his own right.

Distribution. Spain (Málaga province).

Table 1. Host plant use and dietary classification for selected Iberian *Andrena* species. *n*, total number of pollen loads; *N*, number of pollen loads from different localities. Plant taxa: ADO, Adoxaceae; AMA, Amaryllidaceae; API, Apiaceae; ASP, Asparagaceae; AST, Asteraceae; BOR, Boraginaceae; BRA, Brassicaceae; CAM, Campanulaceae; CAP, Caprifoliaceae; CAR, Caryophyllaceae; CIS, Cistaceae; CRA, Crassulaceae; EUP, Euphorbiaceae; FAB, Fabaceae; FAG, Fagaceae; FRA, Frankeniaceae; GER, Geraniaceae; HYP, Hypericaceae; PAP, Papaveraceae; PLA, Plantaginaceae; PLU, Plumbaginaceae; RES, Resedaceae; RHA, Rhamnaceae; ROS, Rosaceae; SAL, Salicaceae; SAP, Sapindaceae; SCR, Scrophulariaceae. Countries: BE, Belgium; BG, Bulgaria; DZ, Algeria; ESP, Spain; FRA, France; IL, Israel; IR, Iran; MA, Morocco; PT, Portugal; SY, Syria; TJ, Tajikistan; TN, Tunisia.

Species	n	N	Origin (and number) of pollen loads	Result of microscopic analysis of pollen grains (% of pollen grains)	Percentage of pure loads of preferred host	Percentage of loads with preferred host	Host range
Aciandrena Warncke	<u> </u>	l				-	
A. fulica Warncke	12	7	ESP (10), PT (2)	BRA 99.6, CIS 0.4	91.7	100.0	Broadly oligolectic (Brassicaceae)
A. vacella Warncke	2	2	ESP (2)	BRA 100.0	100.0	100.0	Broadly oligolectic (Brassicaceae)
aegyptiaca-group							
<i>A. alluaudi</i> Benoist	4	3	MA (2), PT (2)	AST 100.0	100.0	100.0	Broadly oligolectic (Asteraceae; Cichorioideae)
<i>Aenandrena</i> Warncke							
A. aeneiventris Morawitz	15	7	ESP (15)	API 100.0	100.0	100.0	Possibly broadly oligolectic (Apiaceae)
A. hedikae Jäger	22	10	ESP (11), MA (9), PT (2)	API 100.0	100.0	100.0	Possibly broadly oligolectic (Apiaceae)
A. hystrix Schmiedeknecht	9	5	ESP (8), PT (1)	BRA 100.0	100.0	100.0	Broadly oligolectic (Brassicaceae)
Avandrena Warncke							
A. avara Warncke	2	2	ESP (2)	GER 100.0	100.0	100.0	Broadly oligolectic (Geraniaceae)
A. melacana Warncke	6	2	ESP (6)	GER 100.0	100.0	100.0	Broadly oligolectic (Geraniaceae)
A. panurgina De Steffani	11	5	ESP (4), FRA (4), PT (3)	GER 93.6, AST 5.0, BRA 1.4	81.8	100.0	Broadly oligolectic (Geraniaceae)
Blandandrena subgen	nov.				ı		
A. blanda Pérez	27	9	ESP (8), MA (19)	RES 100.0	100.0	100.0	Narrowly oligolectic (<i>Reseda</i> , Resedaceae)
Brachyandrena Pittion	i						
A. colletiformis Morawitz	6	4	ESP (5), PT (1)	API 100.0	100.0	100.0	Possibly broadly oligolectic (Apiaceae)
A. miegiella Dours	5	4	ESP (2), MA (2), TN (1)	API 99.8, AST 0.2	80.0	100.0	Possibly broadly oligolectic (Apiaceae)
Chlorandrena Pérez						•	
A. abrupta Warncke	1	1	PT (1)	AST 100.0	100.0	100.0	Probably broadly oligolectic (Asteraceae; Asteroideae)
A. cinerea Brullé	22	15	ESP (6), FRA (3), PT (12), TN (1)	AST 100.0	100.0	100.0	Broadly oligolectic (Asteraceae; Cichorioideae)
A. curtivalvis Morice	1	1	ESP (1)	AST 100.0	100.0	100.0	Broadly oligolectic (Asteraceae; Cichorioideae)
A. elata Warncke	13	5	ESP (13)	AST 100.0	100.0	100.0	Broadly oligolectic (Asteraceae; Asteroideae)
A. leucolippa Pérez	22	10	ESP (12), FRA (10)	AST 100.0	100.0	100.0	Broadly oligolectic (Asteraceae; Asteroideae)
A. rhenana Stöckhert	8	4	ESP (2), PT (6)	AST 100.0	100.0	100.0	Broadly oligolectic (Asteraceae; Cichorioideae)
A. senecionis Pérez	21	15	ESP (11), FRA (3), MA (3), PT (4)	AST 100.0	100.0	100.0	Broadly oligolectic (Asteraceae; Cichorioideae)
Chrysandrena Hedick	e	1	<u> </u>	<u> </u>	I	1	· · · · · · · · · · · · · · · · · · ·
A. fertoni Pérez	4	3	ESP (4)	AST 100.0	100.0	100.0	Broadly oligolectic (Asteraceae; Cichorioideae)

Condension Warnelse	Species	n	N	Origin (and number) of pollen loads	Result of microscopic analysis of pollen grains (% of pollen grains)	Percentage of pure loads of preferred host	Percentage of loads with preferred host	Host range
A. mandager Peter 10 5 ESP (3), MA (7) BRA 61.8, ROS 22.6, 30.0 80.0 Telyleccic s. st.	Cordandrena Warnek				(%) of ponen grains)	preferred nost	preferred nost	
A reminist		_	5	ESP (3), MA (7)	1	30.0	80.0	Polylectic s. str.
Doublemis Gribodo	Cryptandrena Pittioni			I	I	l		
A. mucida Kriechbaumer (1)		39	10	ESP (29), FRA (10)		76.9	97.4	Polylectic with a strong preference (Fabaceae)
Sriechbaumer (1° generation)	<i>Didonia</i> Gribodo					1	•	
MA (1) Caprifoliacea Capridoliacea Cap	Kriechbaumer (1st	3	3		ASP 100.0	100.0	100.0	Possibly narrowly oligolectic (<i>Muscari</i> ; Asparagaceae)
A. lateranhalder 5	Kriechbaumer (2 nd	12	8		CAP 100.0	100.0	100.0	Broadly oligolectic (Caprifoliaceae)
PF(1) SCR 20.7, PIA 10.3, AST 8.3, CAM 4.7, GER 2.9, CAR 1.2								
A. nebularia S S S ESP (1), MA (4) BRA 100.0 100.0 100.0 Probably broadly of (Brassicaceae)	Pérez		5		SCR 20.7, PLA 10.3, AST 8.3, CAM 4.7,	20.0	20.0	Polylectic s. str.
Warncke			2	ECD (1) MA (4)	DDA 100.0	100.0	100.0	Doct that are the alternation
PT (2) Brassicaceae A Intervalis 7 3 ESP (3), IR (1), TJ (3) API 100.0 100.0 100.0 Broadly oligolectic (A Morrawitz 7 3 ESP (3), IR (1), TJ (3) API 100.0 100.0 100.0 Broadly oligolectic (A Morrawitz 7 3 ESP (3), IR (1), TJ (3) API 100.0 100.0 100.0 Broadly oligolectic (A Lewandrona Hedicke A Leptopyga Pérez 19 12 DZ (1), ESP (1), MA (12), PT (5) BOR 1.8, SCR 0.7 Polylectic with a preference (Resedul, Re A tumetama A fuscua Erichson 18 12 ESP (15), FRA (1), IL (1), PT (1) BRA 52.4, API 15.5, PAP 12.7, AST 9.5, ROS (6.1, EUP 3.1, others 0.7 Polylectic s. st (Rossi) Follows a final part of the standard	Warncke							(Brassicaceae)
Incita-group	A. verticalis Pérez	30	21		BRA 56.3, API 43.8	53.3	53.3	Mesolectic (Apiaceae & Brassicaceae)
Morawitz Leucandrena Hedicke Leucandrena Hedicke July 12 DZ (1), ESP (1), MA (12), PT (5) BOR 1.8, SCR 0.7 Polylectic with a spreference (Reseda, Resola, Resolution) Respectively and the professional preference (Reseda, Resola, Resolution) Respectively and the preference (Reseda, Resolution) Resolution Resolut	incisa-group	l		I				
A. leptopyga Pérez 19 12 DZ (1), ESP (1), MA (12), PT (5) BOR 1.8, SCR 0.7 Polylectic with a s preference (Recela, Re A. tunetama		7	3	ESP (3), IR (1), TJ (3)	API 100.0	100.0	100.0	Broadly oligolectic (Apiaceae)
A. tunetana	<i>Leucandrena</i> Hedicke					1	•	
Schmiedeknecht MA (1) (Brassicaceae) Melanapis Cameron	A. leptopyga Pérez	19		(12), PT (5)		78.9	94.7	Polylectic with a strong preference (<i>Reseda</i> , Resedaceae
A. fiscosa Erichson		4	4		BRA 100.0	100.0	100.0	Probably broadly oligolectic (Brassicaceae)
Melandrena Pérez (1), PT (1) PAP 12.7, AST 9.5, ROS 6.1, EUP 3.1, others 0.7 20.0 86.7 Polylectic s. st 9.5, ROS 6.1, EUP 3.1, others 0.7 A. albopunctata (Rossi) 15 7 ESP (14), MA (1) AST 47.4, API 25.3, CAP 11.2, BRA 6.4, FAB 3.9, PAP 3.1, others 2.7 6.7 73.3 Polylectic s. st 9.5, PAP 3.1, others 2.7 A. assimilis Radoszkowski 15 6 ESP (4), FRA (11) AST 34.8, ROS 21.1, API 21.0, PLU 7.9, AMA 6.6, SAL 2.7, others 6.1 6.7 73.3 Polylectic s. st 9.5, PAP 3.1, API 21.0, PLU 7.9, AMA 6.6, SAL 2.7, others 6.1 A. bicolonata (Rossi) 8 5 ESP (2), PT (6) BRA 100.0 100.0 100.0 Broadly oligolec (Brassicaceae, Papertition of Broadly oligolectic (Arbopana Warncke) 9 8 ESP (3), PT (6) CIS 69.6, API 17.2, AST 10.0, others 3.2 22.2 77.8 Polylectic s. st 10.0, others 3.2 Micrandrena Ashmead A. ampla Warncke 21 10 ESP (10), FRA (10), PT (1) API 100.0 100.0 Broadly oligolectic (Arbopana Probably polylectic (Arbopana Probably polylectic (Arbopana Prezz 2.2) 13 ESP (3), MA (7), PT (10.0) CIS 99.9, FAB 0.1	<i>Melanapis</i> Cameron							
Melandrena Pérez A. albopunctata (Rossi) 15 7 ESP (14), MA (1) AST 47.4, API 25.3, CAP 11.2, BRA 6.4, FAB 3.9, PAP 3.1, others 2.7 20.0 86.7 Polylectic s. st A. assimilis Radoszkowski 15 6 ESP (4), FRA (11) AST 37.48, ROS 21.1, API 21.0, PU 7.9, AMA 6.6, SAL 2.7, others 6.1 6.7 73.3 Polylectic s. st A. bicolorata (Rossi) 8 5 ESP (2), PT (6) BRA 100.0 100.0 100.0 Broadly oligolee (Brassicaceae) (Brassicaceae) A. florentina (Rossi) 9 8 ESP (3), PT (6) CIS 69.6, API 17.2, AST 10.0, others 3.2 22.2 77.8 Polylectic s. st Micrandrena Ashmead A. ampla Warncke 21 10 ESP (10), FRA (10), PT (1) API 100.0 100.0 100.0 Broadly oligolectic (A A. bayona Warncke 2 2 ESP (2) API 50.0, BRA 50.0 50.0 50.0 Probably polylee A. djelfensis Pérez 20 13 ESP (8), FRA (1), MA CIST 99.9, FAB 0.1 95.0 100.0 Broadly oligolectic (C A. fabrella Pérez 22 13 ESP (8), FRA (1)	A. fuscosa Erichson	18	12		PAP 12.7, AST 9.5, ROS	50.0	72.2	Polylectic s. str.
CAP 11.2, BRA 6.4, FAB 3.9, PAP 3.1, others 2.7 A. assimilis Radoszkowski S ESP (4), FRA (11) AST 34.8, ROS 21.1, API 21.0, PLU 7.9, AMA 6.6, SAL 2.7, others 6.1 S ESP (2), PT (6) BRA 100.0 100.0 100.0 Broadly oligolectic (Assimation of the state of the	Melandrena Pérez			I	I			
Radoszkowski	•	15	7	ESP (14), MA (1)	CAP 11.2, BRA 6.4, FAB 3.9, PAP 3.1,	20.0	86.7	Polylectic s. str.
(Rossi) A. florentina Magretti A. morio Brullé (including A. hispania Warncke) Micrandrena Ashmead A. ampla Warncke A. bayona Warncke A. bayona Warncke A. djelfensis Pérez Day Service Day Ser		15	6	ESP (4), FRA (11)	API 21.0, PLU 7.9, AMA 6.6, SAL 2.7,	6.7	73.3	Polylectic s. str.
Magretti (Brassicaceae) A. morio Brullé (including A. hispania Warncke) 9 8 ESP (3), PT (6) CIS 69.6, API 17.2, AST 10.0, others 3.2 22.2 77.8 Polylectic s. st. 10.0, others 3.2 Micrandrena Ashmead A. ampla Warncke 21 10 ESP (10), FRA (10), PT (1) API 100.0 100.0 100.0 Broadly oligolectic (API 100.0) A. bayona Warncke 2 2 ESP (2) API 50.0, BRA 50.0 50.0 50.0 Probably polylectic (API 100.0) A. djelfensis Pérez 20 13 ESP (3), MA (7), PT (10) CIS 99.9, FAB 0.1 95.0 100.0 Broadly oligolectic (COI 100) A. fabrella Pérez 22 13 ESP (8), FRA (1), MA CIST 99.9, AST 0.1 90.0 100.0 Broadly oligolectic (COI 100.0)		8	5	ESP (2), PT (6)	BRA 100.0	100.0	100.0	Broadly oligolectic (Brassicaceae)
(including A. hispania Warncke) 10.0, others 3.2 Micrandrena Ashmead A. ampla Warncke 21 10 ESP (10), FRA (10), PT (1) API 100.0 100.0 100.0 Broadly oligolectic (API of the probably polylectic (API of the probably p		9	5	MA (1), PT (8)	BRA 100.0	100.0	100.0	Broadly oligolectic (Brassicaceae)
Micrandrena Ashmead A. ampla Warncke 21 10 ESP (10), FRA (10), PT (1) API 100.0 100.0 100.0 Broadly oligolectic (API 100.0) A. bayona Warncke 2 2 ESP (2) API 50.0, BRA 50.0 50.0 50.0 Probably polylectic (Control of the probably polylectic) A. djelfensis Pérez 20 13 ESP (3), MA (7), PT (10) CIS 99.9, FAB 0.1 95.0 100.0 Broadly oligolectic (Control of the probably polylectic) A. fabrella Pérez 22 13 ESP (8), FRA (1), MA CIST 99.9, AST 0.1 90.0 100.0 Broadly oligolectic (Control of the probably polylectic)	(including A .	9	8	ESP (3), PT (6)		22.2	77.8	Polylectic s. str.
A. bayona 2 2 ESP (2) API 50.0, BRA 50.0 50.0 50.0 Probably polyled Warncke A. djelfensis Pérez 20 13 ESP (3), MA (7), PT (10) CIS 99.9, FAB 0.1 95.0 100.0 Broadly oligolectic (CIS 99.9, AST 0.1) A. fabrella Pérez 22 13 ESP (8), FRA (1), MA CIST 99.9, AST 0.1 90.0 100.0 Broadly oligolectic (CIS 99.9, AST 0.1)		<u>. </u>	1	<u>I</u>	1	I	I	<u> </u>
Warncke 20 13 ESP (3), MA (7), PT (10) CIS 99.9, FAB 0.1 Pinch (10) 95.0 Pinch (10) Broadly oligolectic (Control (10)) A. fabrella Pérez 22 13 ESP (8), FRA (1), MA CIST 99.9, AST 0.1 Pinch (10) 90.0 Pinch (10) Broadly oligolectic (Control (10))	A. ampla Warncke	21	10		API 100.0	100.0	100.0	Broadly oligolectic (Apiaceae)
(10) A. fabrella Pérez 22 13 ESP (8), FRA (1), MA CIST 99.9, AST 0.1 90.0 100.0 Broadly oligolectic (C		2	2	ESP (2)	API 50.0, BRA 50.0	50.0	50.0	Probably polylectic
	A. djelfensis Pérez	20	13		CIS 99.9, FAB 0.1	95.0	100.0	Broadly oligolectic (Cistaceae)
(5), P1 (8)	A. fabrella Pérez	22	13	ESP (8), FRA (1), MA (5), PT (8)	CIST 99.9, AST 0.1	90.0	100.0	Broadly oligolectic (Cistaceae)

Species	n	N	Origin (and number) of pollen loads	Result of microscopic analysis of pollen grains	Percentage of pure loads of	Percentage of loads with	Host range
4			ECD (O)	(% of pollen grains)	preferred host	preferred host	D11 ·
A. icterina Warncke	9	6	ESP (9)	BRA 69.1, CIS 10.6, SAL 10.3, EUP 5.8, others 4.2	22.2	88.9	Polylectic s. str.
A. longibarbis Pérez	13	8	ESP (2), MA (8), PT (3)	BRA 99.6, AST 0.4	92.3	100.0	Broadly oligolectic (Brassicaceae)
A. nana (Kirby)	51	28	ESP (33), FRA (1), MA (8), PT (9)	API 71.2, BRA 28.6, EUP 0.2	68.6	74.5	Polylectic with a strong preference (Apiaceae)
A. nitidula Pérez	39	18	ESP (16), MA (21), PT (2)	BRA 100.0	100.0	100.0	Broadly oligolectic (Brassicaceae)
<i>A. omnilaevis</i> Wood	6	5	ESP (2), PT (4)	CRA 100.0	100.0	100.0	Probably narrowly oligolectic (<i>Sedum</i> , Crassulaceae)
A. orana Warncke	17	3	DZ (9), MA (5), PT (3)	BRA 100.0	100.0	100.0	Broadly oligolectic (Brassicaceae)
<i>A. pauxilla</i> Stöckhert	11	5	ESP (11)	CRA 82.1, BRA 17.9	81.8	81.8	Possibly polylectic with a strong preference (<i>Sedum</i> , Crassulaceae)
A. spreta Pérez	19	11	ESP (15), MA (3), PT (1)	BRA 93.1, AST 3.0, EUP 2.8, FAB 1.2	78.9	100.0	Polylectic with a strong preference (Brassicaceae)
A. tenuistriata Pérez	39	27	ESP (17), FRA (3), MA (6), PT (13)	BRA 99.8, others 0.2	94.9	100.0	Broadly oligolectic (Brassicaceae)
Nobandrena Warncke			707 (10)				
A. funerea Warncke	12	6	ESP (12)	BRA 100.0	100.0	100.0	Broadly oligolectic (Brassicaceae)
Notandrena Pérez	125		D7 (2) ECD (2) 3 (4)	DD 4 100 0	100.0	100.0	D 11 1: 1 :
A. aerinifrons Dours	25	8	DZ (3), ESP (3), MA (11), PT (8)	BRA 100.0	100.0	100.0	Broadly oligolectic (Brassicaceae)
A. bellidis Pérez	3	3	ESP (2), PT (1)	AST 38.9, RES 36.4, RAN 18.2, BOR 6.5	0.0	66.7	Polylectic s. str.
A. juliana Wood	35	2	ESP (35)	API 82.6, FRA 13.1, CIS 2.8, others 1.4	74.3	82.9	Polylectic with a strong preference (Apiaceae)
A. leucophaea Lepeletier	2	2	ESP (2)	AST 100.0	100.0	100.0	Possibly oligolectic (Asteraceae Asteroideae)
A. nigroviridula Dours	9	8	ESP (4), MA (4), PT (1)	BRA 100.0	100.0	100.0	Broadly oligolectic (Brassicaceae)
A. varuga Warncke	3	2	ESP (3)	BRA 100.0	100.0	100.0	Probably broadly oligolectic (Brassicaceae)
numida-group	Ι.	7	ECD (/) ED (1)	DDA C/O ADIO/C	F0.0	(2.5	3.6 1 . /4 . 0-
A. hypopolia Schmiedeknecht	8	7	ESP (4), FRA (1), PT (3)	BRA 64.8, API 34.6, AST 0.6	50.0	62.5	Mesolectic (Apiaceae & Brassicaceae)
A. ranunculorum Morawitz	17	17	FRA (17)	BRA 75.8, API 10.2, ROS 5.7, FAG 5.6, AST 2.1, ADO 0.5	64.7	100.0	Polylectic with a strong preference (Brassicaceae)
A. monilia	2	2	ESP (1), MA (1)	BRA 100.0	100.0	100.0	Probably broadly oligolectic
Warncke							(Brassicaceae)
Ovandrena subgen. no A. farinosa Pérez	ov. 9	5	ESP (9)	FAB 100.0	100.0	100.0	Broadly oligolectic (Fabaceae)
A. oviventris Pérez	28	11	ESP (9) ESP (9), FRA (4), MA (12), PT (3)	RES 98.3, others 1.7	82.1	100.0	Narrowly oligolectic (<i>Reseda</i> , Resedaceae)
Plastandrena Hedicke	<u></u>		(12),11(0)		<u> </u>	<u> </u>	1 (cocinicac)
A. asperrima Pérez	56	29	ESP (8), FRA (5), MA (43)	BRA 77.6, RES 18.6, ROS 2.5, AST 1.3	69.6	87.5	Polylectic with a strong preference (Brassicaceae)
A. pilipes Fabricius s. str.	28	21	ESP (15), FRA (8), PT (5)	BRA 54.4, ROS 18.9, AST 13.8, CIS 6.6, API 5.7, others 0.7	35.7	57.1	Polylectic s. str.
<i>relata-</i> group	1	1	1	·	I	ı	1
A. corax Warncke	10	4	ESP (8), PT (2)	RES 99.7, AST 0.3	90.0	100.0	Narrowly oligolectic (<i>Reseda</i> , Resedaceae)
A. laurivora Warncke	3	1	MA (1)	RES 100.0	100.0	100.0	Probably narrowly oligolectic (<i>Reseda</i> , Resedaceae)

Species	n	N	Origin (and number) of pollen loads	Result of microscopic analysis of pollen grains (% of pollen grains)	Percentage of pure loads of preferred host	Percentage of loads with preferred host	Host range
A. relata Warncke	2	2	ESP (2)	RES 100.0	100.0	100.0	Probably narrowly oligolectic (<i>Reseda</i> , Resedaceae)
<i>Rufandrena</i> Warncke							
A. orbitalis Morawitz	9	6	ESP (2), FRA (4), PT (3)	PLA 100.0	100.0	100.0	Narrowly oligolectic (<i>Plantago</i> , Plantaginaceae)
A. rufiventris Lepeletier	3	1	MA (3)	PLA 100.0	100.0	100.0	Narrowly oligolectic (<i>Plantago</i> , Plantaginaceae)
Simandrena Pérez							
A. antigana Pérez	25	14	ESP (6), MA (7), PT (12)	BRA 99.8, others 0.2	96.0	100.0	Broadly oligolectic (Brassicaceae)
A. cilissaeformis Pérez	5	5	ESP (2), MA (3)	BRA 83.6, EUP 8.8, RHA 7.6	60.0	80.0	Probably polylectic with a strong preference (Brassicaceae)
A. propinqua Schenck	43	31	BE (4), ESP (21), FRA (5), MA (3), PT (10)	BRA 46.1, ROS 25.8, FAB 10.2, CIS 4.4, CRA 3.2, BOR 2.5, others 7.7	30.2	58.1	Polylectic s. str.
A. rhypara Pérez	4	3	MA (4)	RES 100.0	100.0	100.0	Possibly narrowly oligolectic (<i>Reseda</i> ; Resedaceae)
A. vetula Lepeletier	31	16	ESP (20), FRA (2), MA (7), SY (1), TN (1)	BRA 99.8, others 0.2	93.5	100.0	Broadly oligolectic (Brassicaceae)
Truncandrena Warncl	кe						
A. doursana Dufour	8	3	MA (7), PT (1)	BRA 100.0	100.0	100.0	Broadly oligolectic (Brassicaceae)
A. ferrugineicrus Dours	28	18	DZ (1), ESP (16), MA (2), PT (9)	BRA 100.0	100.0	100.0	Broadly oligolectic (Brassicaceae)
<i>A. nigropilosa</i> Warncke	23	8	ESP (16), FRA (5), MA (2)	BRA 100.0	100.0	100.0	Broadly oligolectic (Brassicaceae)
A. villipes Pérez	6	2	FRA (1), PT (5)	CIS 100.0	100.0	100.0	Probably broadly oligolectic (Cistaceae)

Description of missing sexes

Andrena (Micrandrena) alma Warncke, 1975

Description. Male. Body length 6.5–7 mm (Fig. 53A). *Head*: Dark, 1.2 times wider than long. Clypeus flattened, unevenly punctate with large punctures, punctures separated by 0.5–2 puncture diameters, underlying surface shagreened in basal half, polished and shiny in apical half. Process of labrum trapezoidal, slightly wider than long, apical margin slightly thickened. Gena 1.3 times width of compound eye (Fig. 53B, C); ocelloccipital distance 0.5 times diameter of lateral ocellus. Face medially with whitish hairs, scape with mixture of black and white hairs, inner margin of compound eyes with black hairs, gena ventrally with white hairs, becoming black dorsolaterally, vertex and frons with mixture of black and pale hairs, none exceeding length of scape. Antennae dark, A3 exceeding length of A4, shorter than A4+5. *Mesosoma*: Scutum and scutellum finely granularly shagreened and weakly shiny, shallowly and obscurely punctate, punctures separated by 2-3 puncture diameters. Pronotum with clear humeral angle. Mesepisternum and dorsolateral surfaces of propodeum with fine granular microreticulation, with regular slightly raised hair bearing punctures. Propodeal triangle with regular granular shagreen, basally and medially with obscure and finely raised rugosity, propodeal triangle thus defined by change in surface sculpture compared to dorsolat-

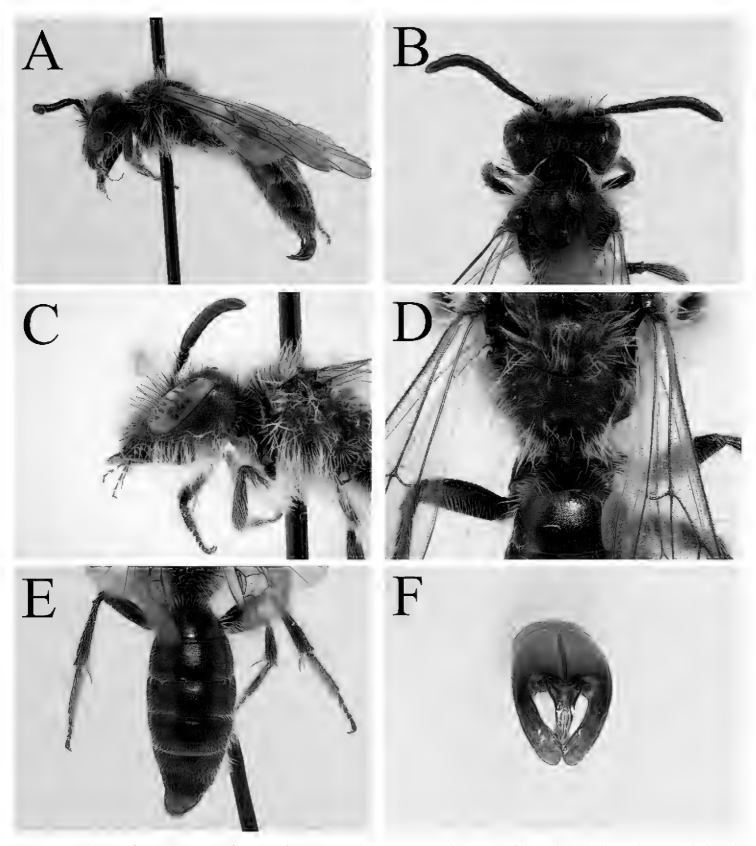


Figure 53. Andrena (Micrandrena) alma Warncke, 1975 male **A** profile **B** head, dorsal view **C** head, profile view **D** propodeal triangle, dorsal view **E** terga, dorsal view **F** genital capsule, dorsal view.

eral parts of propodeum (Fig. 53D). Mesosoma laterally with long white pubescence, exceeding length of scape, pubescence becoming brownish and shorter dorsally, not exceeding length of scape. Legs dark, pubescence whitish. Hind tarsal claws with inner tooth. Wings hyaline, venation and stigma dark brown, nervulus interstitial. *Metasoma*: Terga dark, marginal areas obscurely lightened dark hyaline brown apically (Fig. 53E). Tergal discs with obscure and weak punctures that disappear into even underlying microreticulation, surface weakly shiny. Tergal discs with sparse long light brown hairs, T2–4 laterally with weak, widely interrupted apical hair fringes. S8 columnar, slightly

broadened apically, ventrally covered with short yellowish hairs. Genital capsule compact, gonocoxae apically weakly produced into short rounded teeth; gonostyli parallel-sided, spatulate; penis valves slightly broadened basally (Fig. 53F).

Diagnosis. Andrena alma can be recognised due to its combination of small body size, dark integument, pronotum with humeral angle, evenly shagreened and weakly shiny terga, gena exceeding the width of the compound eye (Figs 53B, C), flattened and striation-free clypeus, simple genital capsule (Fig. 53F), and smooth and granulate propodeal triangle with obscure and finely raised rugosity basally (Fig. 53D). This smooth propodeal triangle places it close to former Distandrena species, but the flattened and striation-free clypeus excludes A. longibarbis Pérez, 1895 (clypeus domed, obscurely striate) and A. orana Warncke, 1974 (clypeus domed and striate). The evenly shagreened terga and obscure basal rugosity on the largely granularly shagreened propodeal triangle place it superficially close to A. djelfensis, but this species can easily be separated by the distinctive genital capsule with elongate and strongly medially bent gonostyli, whereas the genital capsule is simple and lacking distinctive features in A. alma. Finally, A. alma has a distinctly broadened gena that slightly but distinctly wider than the width of the compound eye, an unusual character in *Micrandrena*. This allows separation from A. abjecta which has the gena equalling the width of the compound eye. Collectively, these characters make recognition of A. alma straightforward, though as the males of A. tenostra and A. aff mica are unknown, diagnosis may become more challenging in the future. A level of caution should therefore be applied when identifying material from the extreme south and south-east of Spain.

Distribution. Central and southern Portugal and Spain.

Material examined. Portugal: Algarve, Monte Gordo, Retur, Praia do Cabeço, 29.iii.2022, 1♂, leg. T.J. Wood, TJWC; Algarve, Tavira, Cacela Velha, 28.iii.2022, 1♂, leg. T.J. Wood, TJWC; Spain: Almodóvar del Campo (Ciudad Real), 700 m, 24.iii.2005, 1♂, leg. F.J. Ortiz-Sánchez, FJOS; Santa Ana la Real, Sierra Aracena (Huelva), 630 m, 13.iv.2006, 2♂, leg. F.J. Ortiz-Sánchez, FJOS; El Hongo (P.N. Doñana), 30.iii.2018, 1♂, leg. F. Molina, EBDC.

Andrena (?Euandrena) ramosa Wood, 2022

Description. Male. Body length 8–10 mm (Fig. 54A). *Head:* Dark, 1.05 times wider than long (Fig. 54C). Clypeus long, weakly domed, unevenly punctate, punctures separated by 0.5–2 puncture diameters, underlying surface strongly shagreened to microreticulate in basal half, becoming smooth and shiny in apical half. Process of labrum trapezoidal, 3 times wider than long, ventral surface smooth and polished. Gena equalling width of compound eye; ocelloccipital distance 1.5 times diameter of lateral ocellus. Face medially and gena ventrally with long yellowish hairs, face laterally, frons, and scape with long black hairs, mixing medially on face with yellowish hairs, longest exceeding length of scape. Antennae dark, A3 exceeding length of A4, shorter than A4+5, A4 slightly longer than wide, A5–13 elongate, clearly longer than wide. *Mesosoma:* Scutum and scutellum obscurely punctate, punctures separated by 1–2 puncture di-

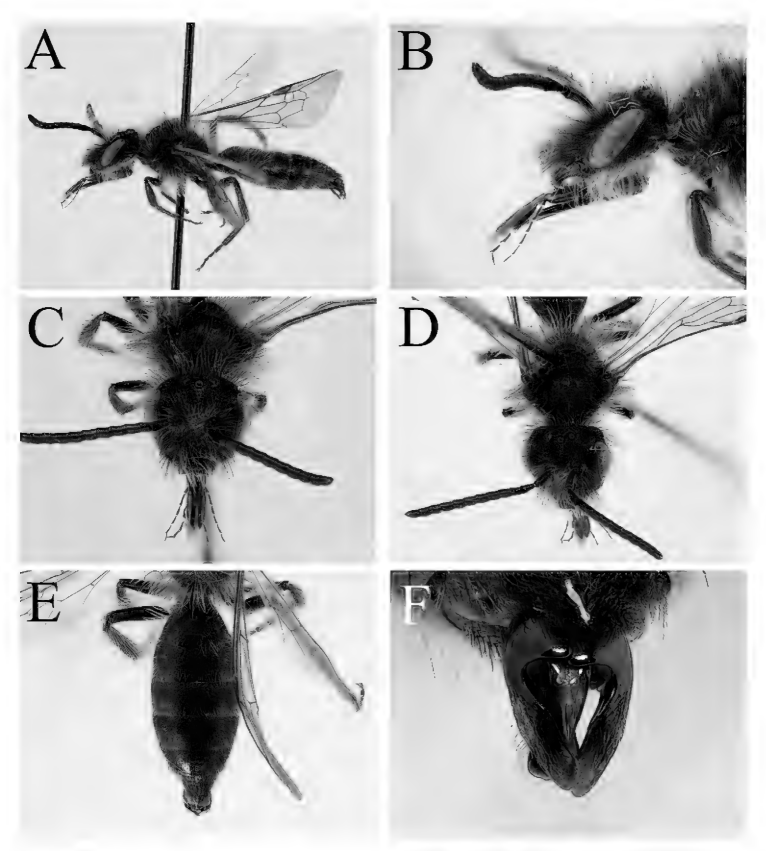


Figure 54. Andrena (?Euandrena) ramosa Wood, 2022 male **A** profile **B** head, profile view **C** head, frontal view **D** scutum, dorsal view **E** terga, dorsal view **F** genital capsule.

ameters, disappearing into extremely strong underlying granular microreticulation, surface dull (Fig. 54D). Pronotum rounded. Mesepisternum and dorsolateral surface of propodeum with fine granular microreticulation, with finely raised network of reticulation that gives impression of large shallow punctures. Propodeal triangle narrow, surface with fine granular shagreen, basally and medially with finely raised rugosity, propodeal triangle thus defined by change in surface sculpture compared to dorsolateral parts of propodeum. Mesosoma with long, densely branches and plumose yellowish hairs clearly exceeding length of scape, black plumose hairs intermixed around wing

bases, on scutum, and propodeum. Legs dark, pubescence brownish to black. Hind tarsal claws with inner tooth. Wings hyaline, stigma dark brown, venation dark to light brown, nervulus weakly antefurcal. *Metasoma*: Terga dark, apical rim of marginal areas very narrowly lightened hyaline brown (Fig. 54E). Tergal discs with obscure hair-bearing punctures, disappearing into background sculpture, becoming more strongly defined laterally, underlying surface shagreened and weakly shiny. T1–3 with long but loose plumose yellowish-brown hairs, these becoming black on T4–5. S8 relatively short, rectangular, apically truncate, ventrally covered with dark brown hairs. Genital capsule moderately elongate, gonocoxae apically weakly produced into short rounded projections, gonostyli more or less parallel-sided, spatulate (Fig. 54F). Penis valves occupying ½ space between gonostyli, slightly narrowing towards their apexes.

Diagnosis. The male of *A. ramosa* is morphologically most similar to *A. (Euandrena) solenopalpa* due to the long head (only marginally wider than long) and clypeus that is shiny at least in its apical half. The two species are easily separated by the mouthparts, as in *A. ramosa* the mouthparts that protrude in front of the head are at most as long as the head (viewed frontally or laterally, Fig. 54B), whereas in *A. solenopalpa* the mouthparts are extremely long, the labial palpi alone exceed the length of their head, the part of the mouthparts protruding in front of the head therefore collectively greatly exceed the length of the head. The clypeus of *A. solenopalpa* is also more extensively shiny, the gonostyli have their outer margin emarginate and are apically produced into narrow points, and A3 slightly exceeds the length of A4+5, whereas in *A. ramosa* the clypeus is shiny only in its apical half, the gonostyli are spatulate and apically truncate, and A3 only slightly exceeds A4, and is shorter than A4+5. The two species do not occur in sympatry, with *A. ramosa* restricted to south-western Spain, whereas *A. solenopalpa* is found in central and eastern Spain to southern France.

Distribution. South-western Spain (Cádiz, Sevilla).

Remarks. The phylogenetic placement of A. ramosa remains somewhat obscure even following the discovery of the male sex and generation of a barcode sequence. A 658-bp fragment was generated from the female type specimen [BOLD accession] number: IBIHM524-21], but this did not fall unambiguously close to any species or subgenus. The most similar sequences belonged to the subgenus Euandrena, specifically to A. symphyti (90.26%), A. montana Warncke, 1973 (90.31%), A. fulvida Schenck, 1853 (89.98%), and *A. rufula* Schmiedeknecht, 1883 (89.84%). Morphologically, A. ramosa does not fall nicely into Euandrena, as the female sex has foveae which are narrow but which do not narrow ventrally. However, *Euandrena* are part of the most highly derived clade of Andrena (Pisanty et al. 2022b), and subgeneric classification in this clade has been highly problematic due to the lack of clear delineating characters. Andrena ramosa does not belong to Margandrena Warncke, 1968 due to the lack of a strong humeral angle on the pronotum. It does not belong to the *crocusella*-group due to the lack of a humeral angle and the lack of lateral projections on the male penis valves (in addition to the lack of barcode similarity), and whilst it has strongly plumose pollen collecting hairs comprising the propodeal corbiculae and femoral scopae, those of the tibial scopae are simple, and the foveae do not narrow below which makes

placement in *Chrysandrena* Hedicke, 1933 difficult. *Andrena ramosa* is therefore best considered to be affiliated with the subgenus *Euandrena*, pending investigation with more powerful genetic techniques. It clearly possesses a unique and unusual morphology within the West Palaearctic *Andrena* fauna.

Examination of additional material from the province of Cádiz has shown that *A. ramosa* is commonly encountered in the Parque Natural Los Alcornocales area. Here it can be encountered between January and March, and is most frequently observed on *Erica* (Ericaceae; Pérez Gómez in litt.). However, the pollen host is still obscure, since none of these bees have been observed collecting pollen. Moreover, Ericaceae pollen is small, with the grains typically having a diameter of 25 µm. The widely spaced and strongly branched and plumose pollen collecting hairs of *A. ramosa* (described and illustrated by Wood et al. 2022a) would not seem to be an adaptation to the collection of small Ericaceae pollen grains, and indeed this adaptation is absent in the Ericaceae specialist *A. (Cnemidandrena) fuscipes* (Kirby, 1802) which has simple pollen collecting hairs. Further study is required; the assumption remains that *A. ramosa* collects pollen from a plant species with large pollen grains, thus necessitating these branched and plumose hairs.

Material examined. Spain: Carretera Marrufo, Herriza (Cádiz; 3 km E Puerto de Gáliz), 11.xi.2020, 1♀, leg. Á. Pérez Gómez, APGC; Sevilla, Los Pinares de Aznalcázar [37.2782°N, -6.2356°E], 10.iii.2020, 1♀, leg. F. Molina, OÖLM (holotype); Cádiz, Sierra de Montecoche, 31.i.2022, 4♂, 1♀, leg. Á. Pérez Gómez, APGC/TJWC; 18.i.2021, 1♂, leg. Á. Pérez Gómez, APGC; Cádiz, Pico del Montero, 2.ii.2022, 3♂, 1♀, leg. Á. Pérez Gómez, APGC/TJWC; Cádiz, Sierra de Fates, 21.iii.2022, 1♀, leg. Á. Pérez Gómez, APGC; Cádiz, Pico del Montero, Alcalá de los Gazules, 26.iii.2022, 1♂, leg. Á. Pérez Gómez, APGC.

Additional designation of lectotypes

Andrena (Aenandrena) hystrix Schmiedeknecht, 1883

Andrena (Aenandrena) hystrix Schmiedeknecht, 1883: 618, ♀ [France, lectotype by present designation: RMNH].

Remarks. Schmiedeknecht (1883) described several *Andrena* species from material that had been sent to him by Pérez. In each case, he indicated this in his title, e.g. 'Andrena hystrix Perez in litt' [sic]. For several of these species, the location of type material has been unclear, as they mostly did not seem to be in the MNHN in the Pérez collection (see Le Divelec 2021), or elsewhere (see Gusenleitner and Schwarz 2002). Searches in the RMNH unexpectedly uncovered specimens of four species described by Schmiedeknecht with labels written in Pérez's distinctive handwriting. The exact providence of these specimens is unclear, but the RMNH collection is known to contain material from Schmiedeknecht's collection (F. Bakker, pers. comm.). A lectotype was designated for one of these species (*A. (Andrena) mitis* Schmiedeknecht, 1883) in a previous publication (Wood 2023a); the others are designated here.

Schmiedeknecht (1883) described *A. hystrix* from female specimens from Hungary (*Hungaria*) and southern France (*Gallia meridionali*). He did not specify a type, though Gusenleitner and Schwarz (2002) list south France as the *locus typicus*. A specimen of *A. hystrix* from Marseille is labelled with Pérez's handwriting, and this is considered to be part of the original syntypic series; it is here designated as a lectotype (Fig. 55A, B). No specimens from Hungary s.l. could be found.

Material examined. France: Marseille [43.3612°N, 5.3942°E], 1♀, RMNH (lectotype by present designation; Fig. 55A, B).

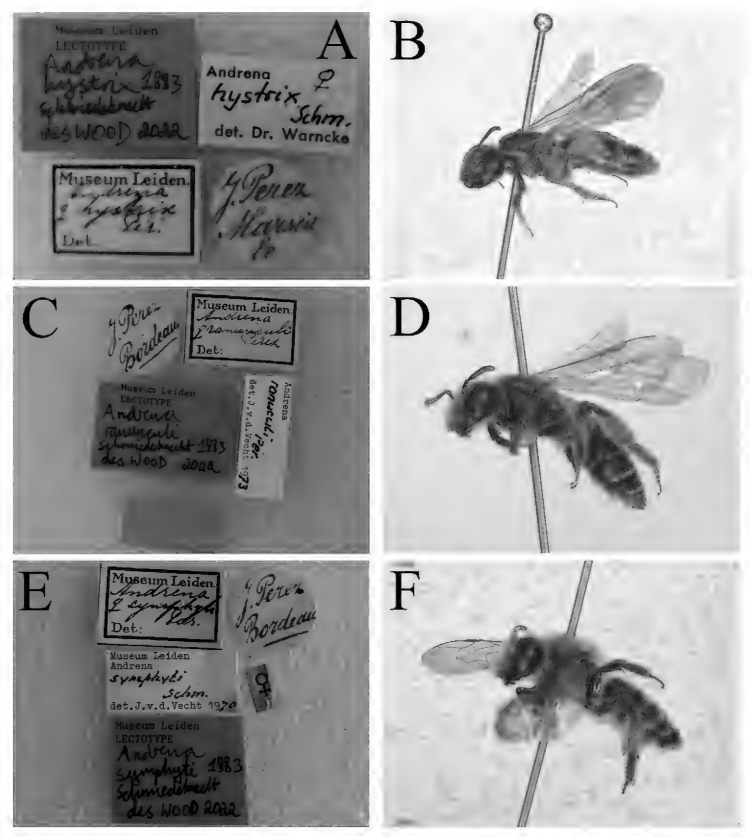


Figure 55. Andrena (Aenandrena) hystrix Schmiedeknecht, 1883, female lectotype **A** label details **B** profile; Andrena (Notandrena) ranunculi Schmiedeknecht, 1883, female lectotype **C** label details **D** profile; Andrena (Euandrena) symphyti Schmiedeknecht, 1883, female lectotype **E** label details **F** profile.

Andrena (Notandrena) ranunculi Schmiedeknecht, 1883

Andrena (Notandrena) ranunculi Schmiedeknecht, 1883: 617, ♀♂ [France, lectotype by present designation: RMNH].

Remarks. Schmiedeknecht (1883) described *A. ranunculi*, comparing it to *A. ranunculorum*. He gave the habitat as Russia (referring to *A. ranunculorum*) and southern France (referring to the specimens received from Pérez). A series of males and females labelled by Pérez as being from Bordeau [sic, = Bordeaux] were found in the RMNH. These conform to Schmiedeknecht's description, and the concept used by subsequent authors. A female is here designated as a lectotype (Fig. 55C, D).

Material examined. France: Bordeaux [44.8352°N, -0.5888°E], 1♀, RMNH (lectotype by present designation; Fig. 55C, D); Bordeaux, 8♂, 5♀, RMNH.

Andrena (Euandrena) symphyti Schmiedeknecht, 1883

Andrena (Euandrena) symphyti Schmiedeknecht, 1883: 583, ♀♂ [France, lectotype by present designation: RMNH].

Remarks. As for the previous two species, material labelled by Pérez was found in the RMNH collection. Two females and one male labelled as being from Bordeau [sic, = Bordeaux]. Schmiedeknecht (1883) specifically states that the type material comes from Bordeaux, writing "In Gallia prope Bordeaux a Dom. Illustrissimo Perez detecta". A female is here designated as a lectotype (Fig. 55E, F).

Material examined. France: Bordeaux [44.8352°N, -0.5888°E], 1♀, RMNH (lectotype by present designation; Fig. 55E, F); Bordeaux, 1♂, 1♀, RMNH.

Designation of neotypes

Andrena (Chlorandrena) boyerella Dours, 1872

Andrena (Chlorandrena) distincta Lucas, 1849 nec. Smith, 1847 [Algeria: MNHN, not examined].

Andrena (Chlorandrena) boyerella Dours, 1872: 429, ♀♂ [Morocco: OÖLM].

Neotype. Morocco: Fès-Meknès, Azrou, 4 km SWW of Bakrit, Cascades Bakrit, 33.0466°N, -5.2681°E, 1650 m, 17.v.2022, 1 \circlearrowleft , leg. T.J. Wood, OÖLM [BOLD accession number WPATW495-22] (Fig. 56).

Remarks. As discussed above, Dours (1872) described *A. boyerella* from southern France and Algeria. No material is available for study, as all of Dours' types were destroyed in a fire. Given that two taxa are present in these regions, in order to fix the name *A. boyerella* on the North African population, a barcoded neotype is designated from Moroccan material (Fig. 56).

Distribution. Morocco, Algeria, Tunisia, Italy (Sicily).

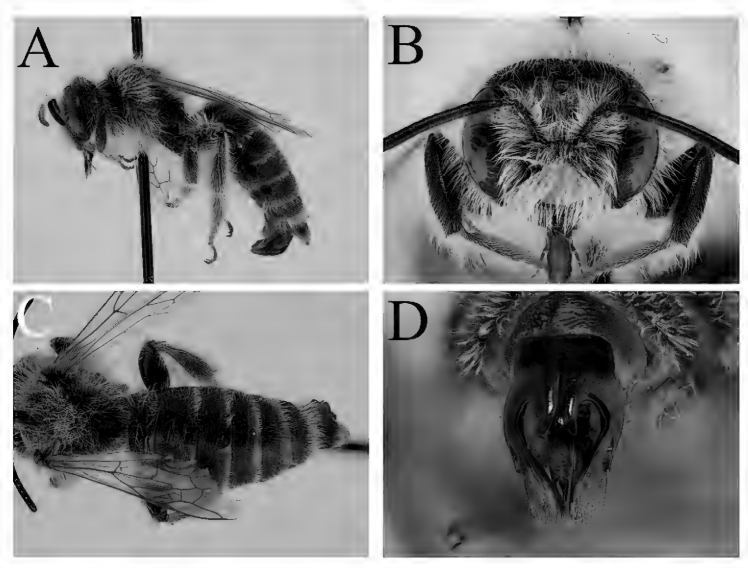


Figure 56. Andrena (Chlorandrena) boyerella Dours, 1872, male neotype **A** profile **B** face, frontal view **C** terga, dorsal view **D** genital capsule dorsal view.

Andrena (Notandrena) griseobalteata Dours, 1872

Andrena (Notandrena) erythrocnemis auctorum. nec. Morawitz, 1871. Andrena (Notandrena) griseobalteata Dours, 1872: 427, ♀ [France: RMNH].

Neotype. France: Pyrénées-Atlantiques, Bérenx [43.4994°N, -0.8575°W], 6.vi.1987, 1\,\text{, leg. E.A.M. Speijer, RMNH (Fig. 57).

Remarks. The correct name to apply to this distinctive taxon has been confused for many years. Through the combination of its large size (for a *Notandrena*) and densely punctate scutum it is comparable only to *A. ungeri* Mavromoustakis, 1952. The name *A. erythrocnemis* Morawitz, 1871 was used by many authors to refer to this taxon (e.g. Warncke 1967), but the lectotype of *A. erythrocnemis* is actually *A. chrysosceles* (see Proshchalykin et al. 2017; Astafurova et al. 2021). Gusenleitner and Schwarz (2002) resolved this issue by resurrecting *A. griseobalteata* to species status. Finally, Wood and Monfared (2022) removed *A. emesiana* Pérez, 1911 (southern Turkey, Syria, Iran) from synonymy with this taxon.

Although Gusenleitner and Schwarz (2002) would seem to have resolved the issue, there is no type specimen for *A. griseobalteata* due to the loss of Dours' collection. In the original description, Dours (1872: 428) listed the species from Saint-Sever in the department of Landes in south-western France, but also from Algeria. This is peculiar,

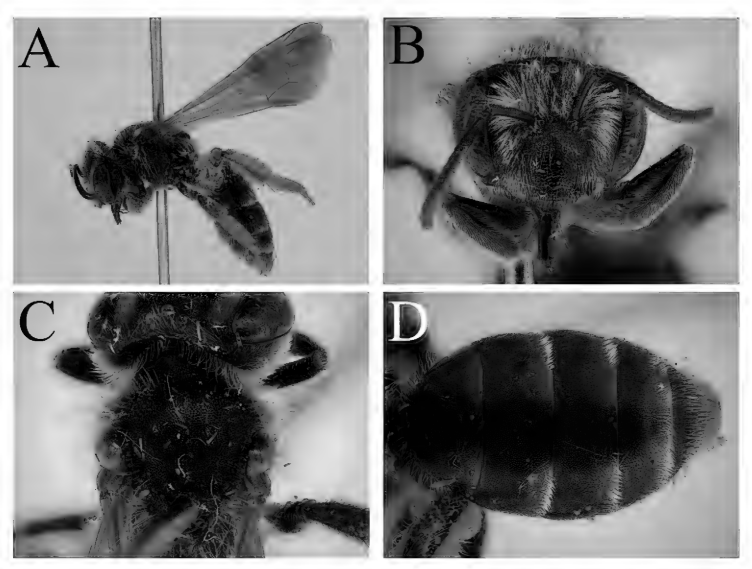


Figure 57. Andrena (Notandrena) griseobalteata Dours, 1872, female neotype **A** profile **B** face, frontal view **C** scutum, dorsal view **D** terga, dorsal view.

because *A. griseobalteata* is not known from North Africa. The original description also does not allow for completely unambiguous recognition of the species, though the dense punctation of the scutum is mentioned. In the interests of nomenclatural stability, a neotype from Bérenx in south-western France (36 km from Saint-Sever) is designated in order to fix the species concept for the future (Fig. 57).

Finally, though listed from Spain by Ortiz-Sánchez (2011, as *A. griseobalteata*; 2020, as *A. erythrocnemis*), the presence of this taxon in Spain is somewhat doubtful. The distribution maps of Warncke (Gusenleitner and Schwarz 2002) show dots around southwestern France into the Pyrenees, but there are no unambiguous dots for Spain. However, I have been able to examine one specimen of *A. griseobalteata* from Spain, from the Sistema Central. The species is also expected to occur in the western Pyrenees; more recent material should be found to establish whether this taxon remains present in Iberia.

Material examined. Spain: Sierra de Gredos, 12 km SSW Hoyos del Espino, 1950–2100 m, 4.vii.1972, 1, leg. J.A.W. Lucas, RMNH.

Distribution. Spain, France, Italy, Croatia, Hungary, Albania, Romania, North Macedonia, Bulgaria, Greece, Turkey (western and northern Turkey; Gusenleitner and Schwarz 2002). The species is not considered to be present in North Africa, or in the Levant, as it is not found in very dry environments.

Andrena (Taeniandrena) poupillieri Dours, 1872

Andrena (Taeniandrena) poupillieri Dours, 1872: 430, ♀ [Algeria: OÖLM].

Andrena (Taeniandrena) poupillieri incana Warncke, 1975a: 310, ♀♂ [Spain, Mallorca: OÖLM, examined].

Neotype. ALGERIA: Tizi-Ouzou, Tigzirt, 36.8877°N, 4.1140°E, 6 m, 31.iii.2017, 1\(\sigma\), leg. H. Ikhlef, OÖLM [BOLD accession number HYMAA322-22] (Fig. 58).

Remarks. This is the taxon referred to as 'A. poupillieri 2' by Praz et al. (2022). The taxon A. poupillieri incana Warncke, 1975 which was described from and restricted to the Balearic Islands is considered a simple synonym of A. poupillieri due to the lack of genetic differentiation observed (see above), even though the tergal punctation is slightly reduced compared to the nominate taxon. The specimen used in the analysis of Praz et al. (2022) is designated as a lectotype (Fig. 58).

Distribution. Morocco, Algeria, Tunisia, Spain (mainland and Balearic Islands). Records (Gusenleitner and Schwarz 2002) from Libya must be confirmed, though they probably do refer to true *A. poupillieri*. Records from Crete probably refer to unrecognised *A. ovata* specimens, and so *A. poupillieri* is not considered to be present there until definitive material is located.

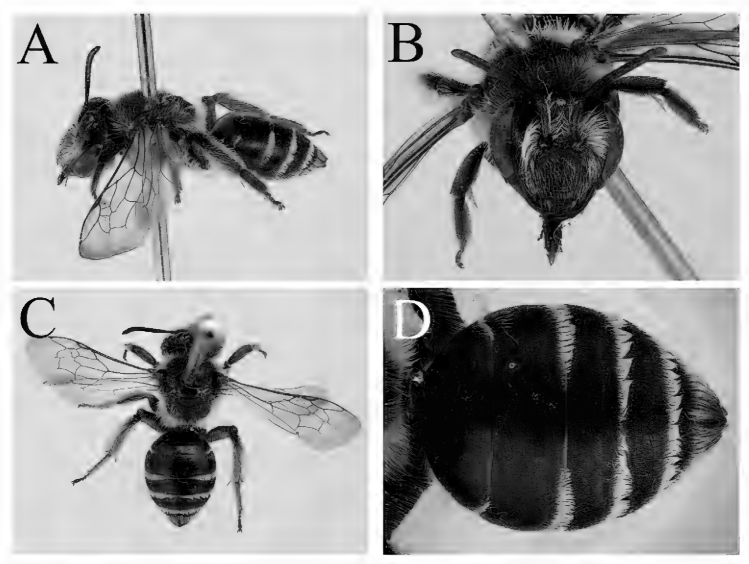


Figure 58. Andrena (Taeniandrena) poupillieri Dours, 1872, female neotype **A** profile **B** face, frontal view **C** dorsal view **D** terga, dorsal view.

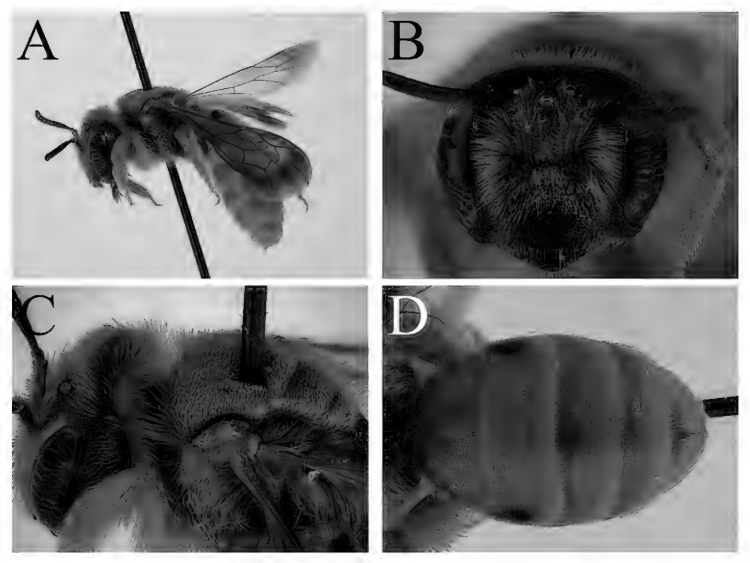


Figure 59. Andrena (Pruinosandrena) succinea Dours, 1872, female neotype **A** profile **B** face, frontal view **C** mesosoma, dorsolateral view **D** terga, dorsal view.

Andrena (Pruinosandrena) succinea Dours, 1872

Andrena succinea Dours, 1872: 424, ♀ [Morocco: OÖLM].

Neotype. Morocco: Oriental, Guercif, P5427, 2 km SW of Bou Rached, 33.8844°N, -3.6154°W, 950 m, 13.v.2022, 1\$\bigsige\$, leg. T.J. Wood, OÖLM [BOLD accession number WPATW389-22] (Fig. 59).

Remarks. As discussed above, it is preferable to designate a neotype for *A. succinea* in order to maintain nomenclatural stability. The barcoded specimen pictured in Fig. 34B is selected as a neotype (Fig. 59) in order to fix the species concept for the future.

Distribution. Morocco, Algeria, Tunisia, Libya, Egypt, Israel and the West Bank, Jordan, Syria, Saudi Arabia, Iran (Wood and Monfared 2022).

Andrena (incertae sedis) numida Lepeletier, 1841

Andrena numida Lepeletier, 1841: 252, ♀ [Morocco: OÖLM].

Neotype. Morocco: Fès-Meknès, Azrou, P7311, 10 km S of Ain Leuh, 1750 m, 33.2220°N, -5.3411°W, 18.v.2022, 1\(\sigma\), leg. T.J. Wood, OÖLM [BOLD accession number WPATW484-22] (Fig. 60).

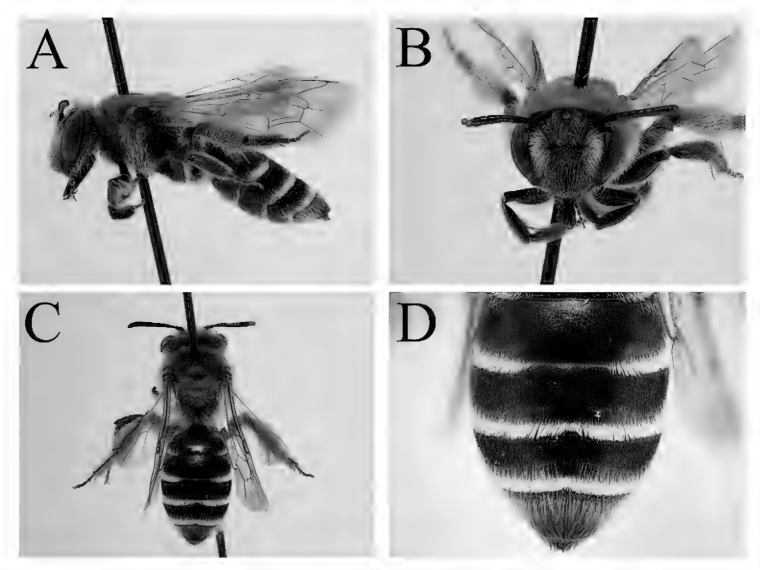


Figure 60. Andrena (incertae sedis) numida Lepeletier, 1841, female neotype **A** profile **B** face, frontal view **C** dorsal view **D** T2–5, dorsal view detail.

Remarks. As discussed above, it is beneficial to designate a neotype for *A. numida* since the original type series cannot be located in the MNHN, and so that the name and genetic identity of North African populations can be fixed.

Distribution. Morocco, Algeria, Tunisia, Libya, Italy (Sicily, Calabria, Campania).

Updated faunal list

Ortiz-Sánchez (2020) lists 212 species of *Andrena* from Spain. However, given the taxonomic and faunal work that has been conducted since then, this total is slightly too low and contains mistakes resulting from taxonomic confusion and the use of variable species concepts in the literature.

Species removed from baseline list

Following the changes detailed in the previous sections, *A. boyerella*, *A. creberrima*, *A. curtula*, *A. hispania*, *A. mariana* s. str., *A. potentillae*, *A. pusilla*, and *A. truncatilabris* are removed from the Iberian fauna following their listing by Ortiz-Sánchez (2020). Four further species must also be removed, *A. similis* Smith, 1849, *A. enslinella* Stöckhert, 1924, *A. similima* Smith, 1851, and *A. chrysopyga* Schenck, 1853. *Andrena* (*Taeniandrena*) *similis*

was shown to be a synonym of *A.* (*Taeniandrena*) russula Lepeletier, 1841 by Praz et al. (2022). Ortiz-Sánchez (2020) listed both species as present.

Dardón (2010) and Dardón et al. (2014) listed A. (Micrandrena) aff enslinella as present in Spain based on the listing of Warncke (1976) and the specimen in his collection. I do not consider this to be ecologically possible, as A. enslinella is a species of Central and Eastern Europe to the Caucasus and Iran, having its western limit in Germany and Austria. The species appears to be absent from France (Le Divelec 2021). Given the highly interesting Andrena fauna in the Sistema Ibérico that contains relictual North African taxa such as A. (Cordandrena) vaulogeri Pérez, 1895 and the endemic A. (Parandrenella) taxana Warncke, 1975, further study is required, as this specimen may represent an undescribed species, as suggested by Dardón (2010).

Andrena (Cnemidandrena) simillima was listed by Ortiz-Sánchez (2020), but not by Warncke (1976). The distribution maps of Warncke (Gusenleitner and Schwarz 2002) give a single isolated dot for this species in north-eastern Spain, presumably in the eastern Pyrenees. The next closest record comes from western France (c. 400 km), with no other records for the Pyrenees. Warncke et al. (1974: carte 140) gave Bordeaux as the south-western limit for A. simillima (specimen leg. Pérez, coll. Warncke OÖLM). I consider the dot on the maps of Gusenleitner and Schwarz (2002) to be of doubtful providence; I have examined no material from Iberia of this species, and I could not locate a Spanish specimen in the Warncke collection. No other authors have ever recorded A. simillima from the Pyrenees, though A. (Cnemidandrena) nigriceps (Kirby, 1802) is common in this region. Males of A. simillima and A. nigriceps are extremely difficult to separate, and it is possible that this dot represents a misidentification of A. nigriceps. Moreover, this record is ecologically questionable, since A. simil*lima* is predominantly a northern species (sensu lato, since cryptic taxa may be present) found in temperate habitats. In the absence of available specimens, the isolated nature of this record, and the identification difficulties associated with this group, I remove A. simillima from the Iberian fauna.

Finally, A. (Melandrena) chrysopyga is listed as present in Iberia. I have seen no Iberian material of this taxon which is generally very rare in collections. It is often confused with forms of A. gravida with light hairs in the terminal fringe. In the distribution maps of Warncke (Gusenleitner and Schwarz 2002), there are dots for A. chrysopyga from south-western France, but none in Iberia proper. Andrena chrysopyga is a species of dry grassland and steppe, extending east across the Great Eurasian Steppe to Kazakhstan. Given the absence or scarcity of steppe habitats in France, its presence in much of the country seems ecologically unlikely. Given the overly generous distribution of A. chrysopyga given by Warncke (e.g. the species is known only from the extreme east of Belgium yet Warncke's map indicates the presence of this taxon throughout Belgium, see Wood 2023a), the lack of available specimens, and the identification mistakes known to occur between A. chrysopyga and A. gravida, A. chrysopyga is not considered to be part of the Iberian fauna until such a time as validated specimens can be found or the species can be newly captured. Its historical and contemporary distribution in France must also be clarified, as I believe that most records are misidentifications of A. gravida.

Species added to baseline list

In addition to the species elevated above or newly described below, the following 15 Andrena species were explicitly added to the Iberian fauna by the following works: A. (incertae sedis) laurivora, A. (Notandrena) juliana Wood, 2021, A. (Euandrena) fortipunctata Wood, 2021, A. (Taeniandrena) benoisti Wood & Praz, 2021, and A. (Taeniandrena) levante Wood & Praz, 2021 (Wood et al. 2021); A. (Plastandrena) nigrospina Thomson, 1872 (Ortiz-Sánchez et al. 2022); A. (Taeniandrena) afzeliella (Kirby, 1802) and A. (Taeniandrena) ovata Schenck, 1853 (Praz et al. 2022); A. (Avandrena) erodiorum Wood & Ortiz-Sánchez, 2022, A. (Avandrena) melacana Warncke, 1967, A. (Taeniandrena) lusitania Wood & Ortiz-Sánchez, 2022, and A. (Suandrena) gades Wood & Ortiz-Sánchez, 2022 (Wood & Ortiz-Sánchez, 2022); A. (incertae sedis) ramosa Wood, 2022 (Wood et al. 2022a); A. (Taeniandrena) contracta Wood, 2022 (Wood 2022); and A. (Andrena) clarkella (Kirby, 1802) (Álvarez Fidalgo and Aguado Martín 2022).

Therefore, relative to the baseline of Ortiz-Sánchez (2020), 12 species are removed, and 28 species are added (including *Andrena* aff *mica*). In sum, the taxonomic changes and faunal updates presented here and in the referenced papers brings the Iberian *Andrena* fauna to 228 species, a substantial increase on the 175 species recorded by Warncke (1975a), with 228 species known from mainland Spain and 128 species from mainland Portugal. A full checklist can be found in Suppl. material 1.

Dietary niche of Iberian Andrena species

Results are presented here for Iberian species for which no or very little previous dietary data have been published. Consequently, these results are not comprehensive, but it is not considered necessary to duplicate here previous analyses that have been conducted in Central Europe (e.g. Westrich 1989) when these same species do not differ in their pollen foraging behaviour in Iberia. Pollen was analysed and removed from a total of 1,127 specimens from 76 species from 12 countries. Findings are summarised in Table 1. *Andrena* species are grouped by subgenus to highlight their often conserved patterns of pollen collection.

Demonstration of oligolecty in understudied species

For many Iberian endemic or West Mediterranean species, an oligolectic dietary niche was clearly and unambiguously demonstrated by pollen analysis. In many cases, specialised pollen use was as expected based on known subgeneric traits, such as the exclusive use of Asteraceae by the subgenus *Chlorandrena* and Brassicaceae by the subgenera *Aciandrena* and *Nobandrena* Warncke, 1968.

It is important to note some pollen collection preferences. Within the *Notandrena*, members of the former *Carandrena* are typically associated with Brassicaceae such as *A. aerinifrons* (Fig. 61A). However, two species deviate from this pattern, *A. bellidis* Pérez, 1895 and *A. leucophaea* Lepeletier, 1841. *Andrena bellidis* appears to be polylectic,

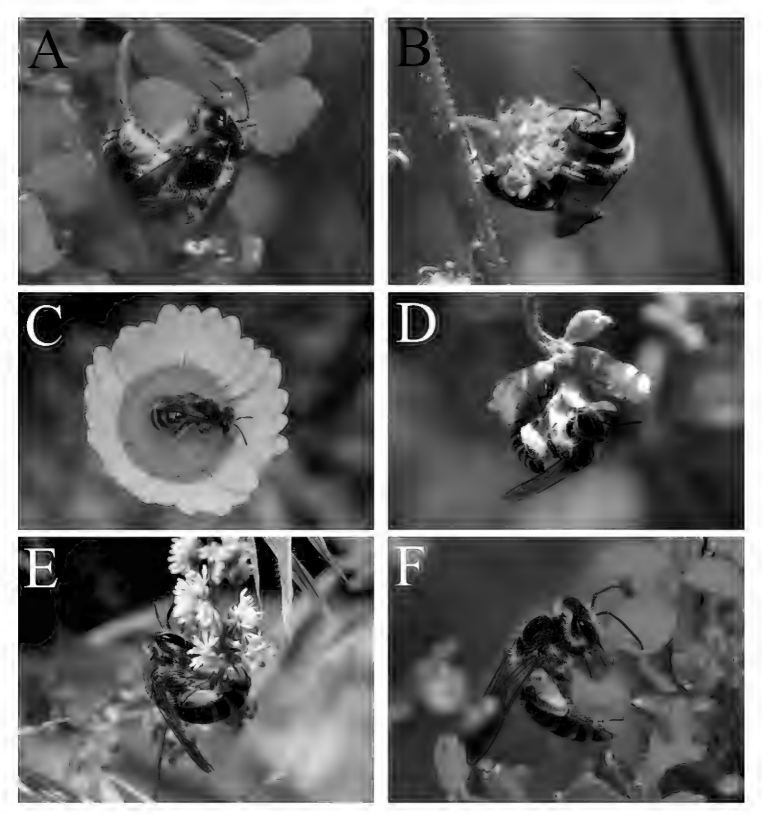


Figure 61. Pollen specialist (oligolectic) Andrena species in Iberia A Andrena (Notandrena) aerinifrons Dours, 1873 (Brassicaceae) B Andrena (incertae sedis) corax Warncke, 1967 (Reseda, Resedaceae) C Andrena (Chlorandrena) elata Warncke, 1975 (Asteroideae, Asteraceae) D Andrena (Ovandrena) farinosa Pérez, 1895 (Lotus dorycnium, Fabaceae) E Andrena (Ovandrena) oviventris Pérez, 1895 (Reseda, Resedaceae) F Andrena (Simandrena) vetula Lepeletier, 1841 (Brassicaceae).

whereas *A. leucophaea* was associated with *Bellis* (Asteraceae; Asteroideae) and may be oligolectic on this subfamily. Both species fly early in the year (typically February-April) and are uncommon in collections; more study is required, but neither species is likely to be specialised on Brassicaceae.

Members of the *relata*-group appear to be specialised on *Reseda* (Resedaceae), including *A. corax* (Fig. 61B). Based on direct observations, all five Iberian members

of the *relata*-group (including *A. macroptera* and *A. murana*) appear to be narrowly oligolectic on *Reseda*, but additional pollen analysis is necessary for confirmation; this being limited by the scarcity of these species in collections.

All studied members of the subgenus *Chlorandrena* are specialists of Asteraceae. However, the only published associations relate to the subfamily Cichorioideae (e.g. Westrich 1989; Schwenninger 2015). However, three Iberian *Chlorandrena* collect exclusively from Asteraceae subfamily Asteroideae: *A. abrupta*, *A. elata* (Fig. 61C), and *A. leucolippa*. These three species form a clade within the *Chlorandrena* along with *A. boyerella* which is also specialised on Asteraceae subfamily Asteroideae (data to be published in the North African revision). This situation resembles the situation within the genus *Panurgus*, members of which are specialised on Asteraceae but for which members of different lineages within *Panurgus* utilise either Asteroideae or Cichorioideae (Wood et al. 2022b).

Within the newly erected subgenus *Ovandrena*, no clear pattern was seen, as *A. farinosa* is a specialist of small-flowered Fabaceae (Fig. 61D) and *A. oviventris* is a specialist of *Reseda* (Fig. 61E). At the moment, *A. farinosa* must be considered a broad oligolege, as the identity of pollen grains removed from museum specimens cannot be identified to the genus level. In Iberia, all my observations of this species come from *Lotus dorycnium* Linnaeus (=*Dorycnium pentaphyllum* Scopoli), and so the species may be more narrowly specialised on the genus *Lotus*, but this requires additional study.

Within the subgenus *Simandrena*, central and northern European species are well-known to be polylectic (Westrich 1989). In contrast, three West Mediterranean species appear to be oligolectic, *A. vetula* (Fig. 61F) and *A. antigana* Pérez, 1895 on Brassicaceae and potentially *A. rhypara* Pérez, 1895 on *Reseda*. All three species belong to the group of *Simandrena* with a strongly shagreened and almost impunctate scutum that is distributed predominantly across the Mediterranean basin, with the highest diversity in North Africa and the Levant. Further study is required to establish the dietary niche of *A. cilissaeformis* which may be polylectic.

Most studied members of the subgenus *Truncandrena* are specialised on Brassicaceae (Westrich 1989), though this subgenus is species-poor in Central Europe and most taxa are Mediterranean and understudied. In Iberia, *A. doursana*, *A. ferrugineicrus*, and *A. nigropilosa* are also specialised on Brassicaceae. However, *A. villipes* was strongly associated with *Cistus* (Cistaceae) based on pollen analysis and direct observations, a behaviour that has not previously been demonstrated for this subgenus. As discussed above, this is likely to be the dietary niche for *A. ghisbaini*, the two species together appear basally in the COI phylogeny (Fig. 29). Additional study is required to establish whether this basal position is supported by more powerful genetic analyses.

The subgenus *Micrandrena* contains mixture of oligolectic and polylectic species in central and northern Europe, though polylectic species predominate (Westrich 1989). In Iberia, numerous species display oligolectic behaviour, specifically *A. ampla* (Apiaceae), *A. djelfensis* and *A. fabrella* Pérez, 1903 (Cistaceae), *A. longibarbis*, *A. nitidula*, *A. orana*, and *A. tenuistriata* (Brassicaceae), and *A. omnilaevis* (Crassulaceae). The dietary status of *A. spreta* is somewhat unclear, as specimens from both the

spring and summer generations analysed here collected predominantly Brassicaceae pollen (93.1%). There is a bias towards spring generation specimens which are typically more abundant, and which fly at a time of year with high Brassicaceae availability. Additional samples from the summer generation are required, but the species can provisionally be considered to be polylectic with a strong preference for Brassicaceae.

There was one species for which the empirical data slightly conflicts with what I believe to be the true dietary niche. For A. (Avandrena) panurgina, the pollen results showed that 93.6% of collected pollen came from Geraniaceae, with the remaining pollen from Asteraceae and Brassicaceae. Following the criteria of Müller and Kuhlmann (2008), this species would not necessarily be classified as oligolectic on Geraniaceae. However, Geraniaceae pollen grains are very large, usually in excess of 50 µm, and it is possible for contaminant pollen to be present between these large grains in a way that is much less common when dealing with smaller grains which pack into scopal hairs more tightly. Behavioural observations of A. panurgina, including those of males which patrol around Erodium (Geraniaceae) plants in March and early April, strongly suggests oligolecty on Geraniaceae. All behavioural pollen-foraging observations of A. panurgina, along with other western Avandrena Warncke, 1968 species A. avara, A. erodiorum, and A. melacana Warncke, 1967 have been made at Erodium (Wood and Ortiz-Sánchez 2022; pers. obs., Álvarez Fidalgo in litt.). Consequently, the presence of Asteraceae and Brassicaceae pollen in the quantitative analysis is considered likely to be contamination, and A. panurgina is considered to be an oligolege of Geraniaceae along with the other western Avandrena species (see Pisanty et al. 2022a).

The data presented here also resolve the pollen collection preferences of the West Mediterranean A. (Rufandrena) orbitalis Morawitz, 1871 and A. (Rufandrena) rufiventris Lepeletier, 1841 which belong to the subgenus Rufandrena Warncke, 1968 that may contain three species, with a further species known from Syria and Hatay province in Turkey (Wood 2023b) which requires genetic confirmation of its placement. The two West Mediterranean species are instantly recognisable in the female sex because of their extraordinarily long and incredibly plumose hairs of the tibial and femoral scopae and both flocculi (see Fig. 64H). Baldock et al. (2018) identified the use of Plantago (Plantaginaceae) pollen by A. orbitalis, but did not come to a firm conclusion as to the overall dietary niche of this species. Analysis of scopal pollen loads and behavioural observations (A. orbitalis in the Algarve, Portugal and A. rufiventris in the Middle Atlas, Morocco) show that both A. orbitalis and A. rufiventris are specialists of Plantago. In the spring, usually in the months of March and April, both species can be seen visiting low-growing Plantago species. Upon arriving (Fig. 62C) at a small flower head, they grab onto its side and present the underside of their body so that falling grains will land on their venter and scopae (Fig. 62A). They will then manipulate the anthers using their fore legs and mandibles (Fig. 62D), often directly scraping pollen from the anthers using their mandibles (Fig. 62E). When the flower head is long, they will sometimes work from bottom to top, often vigorously dislodging pollen grains that can form a small but distinctly visible cloud (Fig. 62B). Pollen landing on the body is then groomed into the scopae.

As *Plantago* is wind-pollinated, it does not provide a nectar source. Nectar is therefore collected from other plants such as *Crepis* (Asteraceae), *Malva* (Malvaceae, Álvarez Fidalgo in litt.), or *Reseda* (Fig. 62F). Pollen is packed into the scopae dry, and inspection of freshly caught and pinned specimens shows that pollen rapidly falls out of the scopae with gentle manipulation, for example with an entomological pin. This

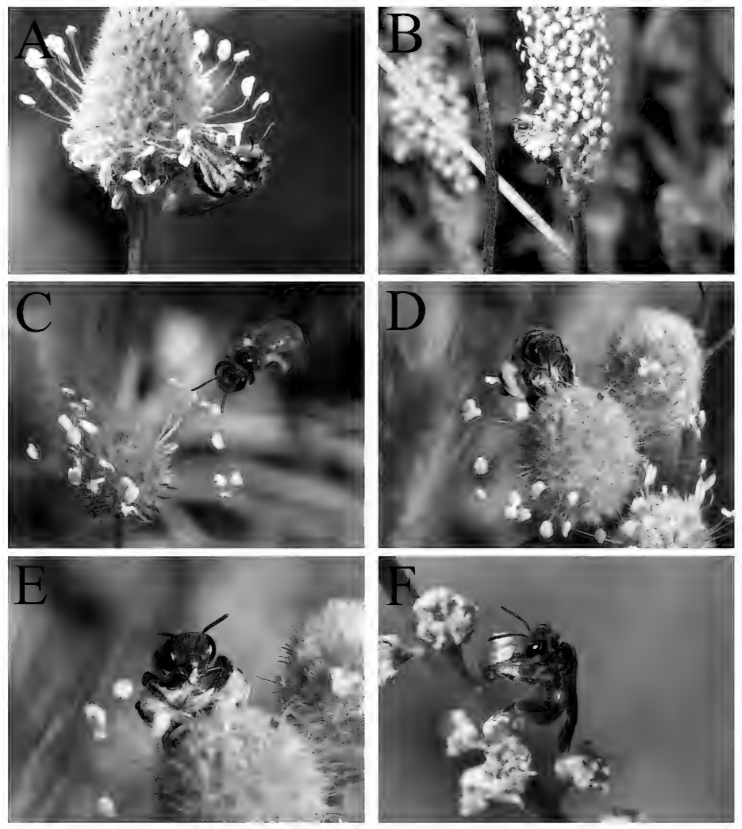


Figure 62. Use of *Plantago* (Plantaginaceae) pollen by *Rufandrena* Warncke, 1968 species. *Andrena* (*Rufandrena*) orbitalis Morawitz, 1871 **A** female, manipulating *Plantago* anther **B** female, vigorously collecting *Plantago* pollen; *Andrena* (*Rufandrena*) rufiventris Lepeletier, 1841 **C** female approaching *Plantago* flower head **D** female manipulating *Plantago* anthers with forelegs and mandibles **E** female scraping pollen from *Plantago* anther using mandibles **F** female drinking nectar at *Reseda* (Resedaceae) with empty scopae.

probably explains both why the scopal hairs are so densely and finely plumose in order to retain these small pollen grains (typically <20 µm in diameter), and also why very few specimens in museum collections have scopae that retain pollen, as dry grains are easily dislodged after specimen collection. Use of pollen from wind-pollinated plants is well-known in bees, predominantly by social polylectic species (Saunders 2018), but also by solitary species, including species belonging to genera such as *Lipotriches* Gerstaecker, 1858 (Halictidae) that can be oligolectic upon members of the Poaceae (Immelman and Eardley 2000). Whilst some *Andrena* species are known to collect pollen from wind-pollinated plants (Wood and Roberts 2017; Saunders 2018), to my knowledge this is the first time that any *Andrena* species has been demonstrated to be specialised for pollen collection on a wind-pollinated plant.

Finally, a note on the pollen collection preferences of A. afzeliella and A. ovatula is beneficial. Praz et al. (2022) demonstrated that A. afzeliella is polylectic with a strong preference for Fabaceae, and A. ovatula is oligolectic on Fabaceae. Additional detail and context can be provided based on observations in Iberia which help to explain their differing distributions and ecologies. Andrena afzeliella typically visits herbaceous Fabaceae, such as *Trifolium pratense*. It visits honestly, and does not manipulate the flowers beyond what is typically expected in an 'honest' visit. It can be found commonly on the open steppe habitats of Old Castile, for example in Segovia, Castroserna de Abajo, Buitreras y Cuevas (25.vii.2021, Fig. 63A, B). In contrast, A. ovatula prefers areas with an Atlantic influence that host members of the tribe Genisteae (Fabaceae), usually woody species (e.g. Cytisus, Genista, Ulex, etc.). Here it visits these flowers honestly for nectar, but roughly manipulates the petals with its two front pairs of legs and its mandibles in order to access the anthers. This can be seen in western and north-western Iberia, as well as in the Sistema Central such as at Segovia, Riofrío de Riaza, Embalse de Riofrío (27. vii.2021, Fig. 63C-F). In my experience, I have never observed A. afzeliella manipulating Fabaceae flowers like this. The relationship with woody Genisteae may explain the earlier emergence of the first generation of *A. ovatula* (peaking April compared to May for A. afzeliella; see typical emergence patterns in north-western Europe in Praz et al. 2022) due to the earlier flowering of woody Fabaceae relative to herbaceous Fabaceae. Consequently, A. ovatula is typically not encountered in open steppe habitats, whereas A. afzeliella typically avoids western parts of Iberia with a strong Atlantic influence.

Pollen use by bivoltine species

A widespread behaviour within *Andrena* is bivoltinism, i.e. species producing a spring and a summer generation. For several bivoltine species, their dietary niche is not yet clear. There is a lack of clarity over two species within *Aenandrena*. For the 15 pollen loads of *A. aeneiventris* (collected between 11th June and 25th July) and the 22 pollen loads of *A. hedikae* (collected between 21st May and 25th July), each load was comprised entirely of Apiaceae pollen. However, both species are bivoltine, and pollen loads from the spring generation must be analysed. Osytshnjuk et al. (2005) state that both species visit Brassicaceae in the spring generation, and Kocourek (1966) gives a

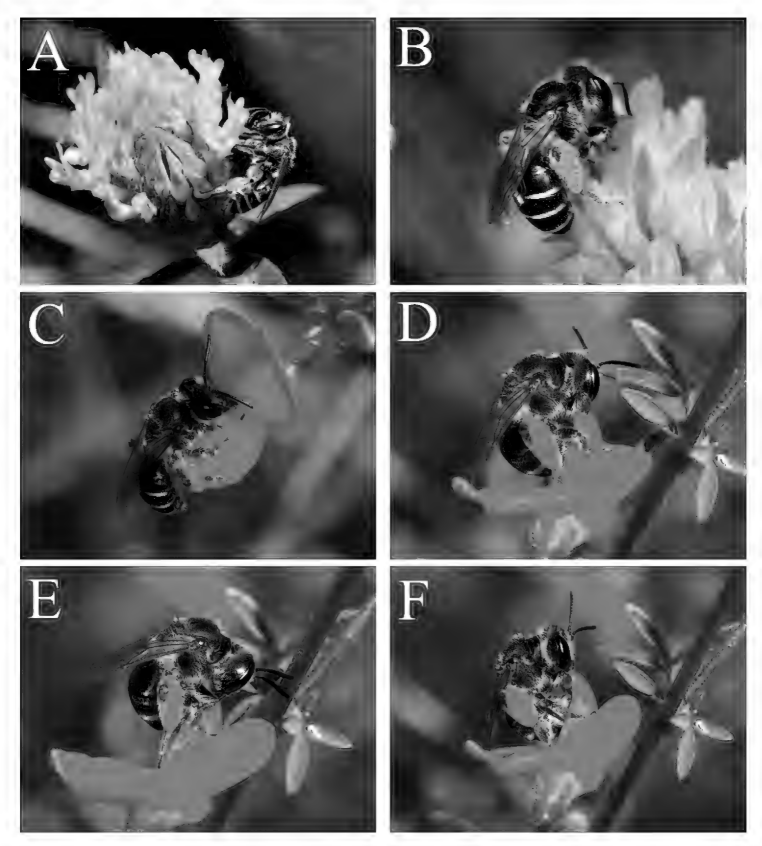


Figure 63. Andrena (Taeniandrena) afzeliella (Kirby, 1802) **A, B** foraging female at Trifolium pratense (Fabaceae); Andrena (Taeniandrena) ovatula (Kirby, 1802) **C-F** foraging female at Genistae spp. (Fabaceae).

list of several Brassicaceae species along with *Sedum* (Crassulaceae) and *Euphorbia* (Euphorbiaceae). I have personally never observed *A. aeneiventris* at flowers of Brassicaceae in the West Mediterranean despite extensively searching this botanical family during the Iberian spring for the large number of univoltine species that specialise on this family. However, in an Iberian context, both species are quite uncommonly observed and collected in their first generation, certainly compared to their abundance in the summer. More study is required, and neither species should be classified as oligolectic on Apiaceae until more pollen load data are available, ideally also from other parts of

southern to south-central Europe. A similar problem is found within the subgenus *Brachyandrena* Pittioni, 1948. Very few pollen loads were available for analysis, but both *A. colletiformis* and *A. miegiella* Dours, 1873 were found to only collect Apiaceae pollen. Both species are also bivoltine. Osytshnjuk et al. (2005) state that *A. colletiformis* visits a variety of botanical families, but prefers the Apiaceae. In this dataset, pollen bearing-specimens of *A. colletiformis* were collected between 25th April and 20th July, thus encompassing both generations. More study is required to establish if pollen sources other than Apiaceae are utilised. Based on observations, they are almost exclusively found on Apiaceae and can be provisionally considered oligolectic on this family.

In contrast, *A. verticalis* provides an example of a bivoltine species that uses Apiaceae, but not exclusively. *Andrena verticalis* flies between March and August, and in this dataset collects 56.3% of its pollen from Brassicaceae (mostly in the spring) and 43.8% of its pollen from Apiaceae (mostly in the summer). However, the use of specific botanical families is not restricted to the two generations, and instead is based on local availability. Interestingly, no mixed pollen loads were detected in the 30 pollen loads examined here, each consisting only of a single botanical family. This recurring pattern of Apiaceae and Brassicaceae use within bivoltine species can be seen for *A. hypopolia*, which also displays mesolecty on these two families. Additional study is required across these aforementioned species to robustly test the limits of mesolecty on Apiaceae and Brassicaceae across two generations (*A. verticalis* and *A. hypopolia*), oligolecty on Apiaceae in both generations (possibly *A. colletiformis* and *A. miegiella*), and possible oligolecty in each generation but upon a different botanical family (potentially *A. aeneiventris* and *A. hedikae*).

Finally, it is necessary to discuss the case of A. mucida which appears to be differentially oligolectic in its spring and summer generation. The terminology surrounding pollen use in bees has been refined and categorised several times in recent years (Cane and Sipes 2006; Müller and Kuhlmann 2008; Kuhlmann and Timmermann 2011; Cane 2020), with debate over how broad or how narrow to make the existing categories. However, to my knowledge, no examples of bee species displaying different specialisation in different generations has been provided or discussed, much less this behaviour formally named. Kocourek (1966) noted the differential foraging behaviour of A. mucida, stating that the 1st generation visited Salix and Muscari, with the 2nd generation visiting Scabiosa. Unlike the case in some bivoltine Andrena species that may display differential levels of specialisation between generations but which cannot unambiguously be assigned to a specific generation morphologically (e.g. some Micrandrena), A. mucida displays simple tibial scopal hairs in the spring generation, and plumose tibial scopal hairs in the summer generation. Individuals can therefore be assigned to each generation with confidence (rather than based only on collection date), and when combined with pollen analysis the data confirm this differential pollen collection behaviour. Therefore, is A. mucida simply a polylectic species, or rather differentially oligolectic? Given the possibly unique nature of this behaviour, it would be beneficial to formally name this pattern of pollen collection heterolecty, in which a species is i) bivoltine, ii) oligolectic in both generations,

and iii) each generation is oligolectic on different botanical families. This name will highlight this highly unusual behaviour, rather than have it hidden away under an all-encompassing 'polylecty'.

Summary for pollen use by Iberian Andrena species

For the 75 Iberian species (not including *A. rufiventris*) for which pollen data are presented, the majority are oligolectic or assessed as likely to be oligolectic. Excluding species for which insufficient data are available to allow confident classification (*A. aeneiventris*, *A. colletiformis*, *A. hedikae*, and *A. miegiella*), 49 species are classified as oligolectic, 21 as polylectic, and one (*A. mucida*) as heterolectic. This high proportion of oligolectic species, most of which have their pollen preferences demonstrated here for the first time, illustrates the degree to which the Iberian fauna i) hosts many specialised species and ii) has been chronically understudied with regard to the basic biology of its constituent bee species. Within the oligolectic species studied here, there is a clear preference for Brassicaceae, this family hosting 22 species, followed by Asteraceae (n=10), Resedaceae (n=6), Cistaceae (n=3), Geraniaceae (n=3), Apiaceae (n=2), and Crassulaceae, Fabaceae, and Plantaginaceae (n=1 each). The broader faunal trends shown by Iberian *Andrena* species will be examined in detail in a subsequent publication.

Identification key

The males of *A. cilissaeformis*, *A. erodiorum*, *A. foeniculae*, *A. juliae*, *A. macroptera*, *A. aff mica*, *A. ortizi*, *A. tenostra*, and *A. urdula* are currently unknown, so they are not included in the key; a level of caution must therefore be taken when working with morphologically similar species. The Spanish *A. allosa* Warncke, 1975 is unknown to me, so the relevant couplet is based on material from the Alps (see Praz et al. 2019).

Unfortunately, after consideration, I have also decided to exclude the male of *A. exigua* from the key because its confident determination is not clear to me. Existing literature (Gusenleitner and Schwarz 2002; Dardón et al. 2014) is light on detail, and my own examination of extremely limited material has not cemented the concept. Due to the nominal rarity of this taxon, no material was available for genetic study, further limiting the confidence with which the sexes can be associated. Confident sex association and determination of *A. exigua* males must remain an open question for students of the Iberian fauna.

For supplementary illustrations, please consult the works cited in the Methodology, particularly Schmid-Egger and Scheuchl (1997) for Euro-Siberian species, as well as my publications describing Iberian species. Moreover, in the near future, the ongoing ORBIT project (providing taxonomic resources for European bees, https://orbit-project.eu/) will provide images for all European *Andrena* species. Between this key and the images presented there, confident determination should be possible in the majority of cases.

The female and male keys are separated for convenience. For the female key, the following shortcuts can be used:

A.	Posterior face of hind femur with latitudinal row of spines or teeth go to 2
B.	Hind tibial spur clearly broadened, either at its base or submedially. Large to
	very large species, at least 12 mm in length
C.	Hairs of the tibial scopae clearly plumose
D.	Propodeal triangle clearly defined by strongly raised carinae, internal surface
	rugose-areolate
E.	Scutum and scutellum with squamous hairsgo to 55
F.	Viewed laterally, propodeal corbicula with internal surface (lateral faces of the
	propodeum) glabrous AND propodeal corbiculae complete (possessing both a
	dorsal and anterior fringe (subgenus Simandrena) go to 64
G.	Fovea strongly constricted medially, strongly diverging from the inner margin of
	the compound eye dorsally (former subgenus <i>Hyperandrena</i>) go to 75
H.	At least some tergal discs extensively red-marked
I.	Head, mesosoma, or tergal discs with metallic reflections (note, A. nigroaenea
	(Kirby) can have bronzy reflections on the terga; if the tibial scopa is composed
	at least partly of orange-red hairs, go to 205)go to 84
J.	Small black species, body length under 8 mm, or if up to 10 mm then with lat-
-	eral faces of the propodeum with clear pattern of raised star-shaped wrinkles (all
	members of the subgenera Aciandrena, Graecandrena, Micrandrena) go to 92
K.	Clypeus flattened over majority of its surface (subgenus <i>Taeniandrena</i>)go to 139
L.	Fovea dorsally narrow, occupying at most 1/3 of space between lateral ocellus and
	compound eye, ventrally narrowing strongly (subgenus <i>Euandrena</i>) go to 152
M.	Clypeus punctate, interspaces forming weakly raised longitudinal wrinkles (for-
	mer subgenus Zonandrena)
N.	Dorsolateral surface of the propodeum reticulate, with large and shallow punctures,
	clearly contrasting the shagreened and shiny propodeal triangle, this lacking lateral
	carinae and becoming shinier on the declivity (subgenus <i>Hoplandrena</i>) go to 168
0.	Pronotum laterally keeled, angulate, keel runs up dorsally to an angled corner.
	go to 174
P .	Large species (over 12 mm in length). Typically with abundant black, brown, and/
	or white pubescence. Clypeus strongly domed. Ocelloccipital distance long, at least
	2 times the diameter of a lateral ocellus (subgenus Melandrena partim)go to 206
Q.	Without this combination of characters; remaining species go to 212
Fema	ales
1	Posterior face of hind femur with latitudinal row of spines or teeth
	(Fig. 64A, B). Note, if there is also a developed latitudinal carina, then the
	spines can be difficult to see (subgenus Lepidandrena, these species with
	squamous hairs on the scutum and scutellum)2
_	Posterior face of hind femur without latitudinal row of spines or teeth. A
	latitudinal carina may be present (e.g. Andrena flavipes)

2	Mesonotum dorsally with short squamous hairs (Fig. 64C, D)	
_	Mesonotum with longer hairs, these never squamous	9
3	At least the disc of T2 red-marked	4
_	All terga dark	5
4	Metasoma entirely red-marked; mesonotum with squamous hairs black	ζ.
	Widespread throughout Iberia, associated with Asphodelus (Asphodelace	-(
	ae)sardoa Lepeletie	r
_	Metasoma with red markings typically restricted to T2-3; mesonotum with	h
	squamous hairs brown-grey. Restricted to the Pyrenees, associated wit	h
	Campanula (Campanulaceae)rufizona Imho	ff
5	T1 comparatively sparsely punctate, large punctures separated by >1 punc	<u></u>
	ture diameter, with scattered micropunctures between these large and rela	l-
	tively coarse punctures. Foveae with outer margin clearly defined, foveae de	3-
	pressed, separated from lateral ocellus by a distance subequal to its diameter	er
	(Fig. 64E)leucolippa Pére	Z
_	T1 finely and densely punctate, punctures separated at most by 1 punc	<u></u>
	ture diameter, usually by 0.5 puncture diameters, without micropunctures	s.
	Foveae with outer margin poorly defined, foveae not clearly depressed	1,
	separated from lateral ocellus by a distance greater than its diamete	r
	(Fig. 64F)	6
6	Galea shagreened and dull, clypeus also shagreened and dull with clea	lľ
	impunctate longitudinal midline. Hind tibiae orange. Associated wit	h
	Ornithogalum (Asparagaceae)baetica Woo	d
_	Galea smooth and shiny, clypeus also at least partially shiny, without im	l -
	punctate longitudinal midline. Hind tibiae dark. Associated with Campanul	a
	(Campanulaceae)	
7	Tarsal segment 5 of the hind leg elongate and strongly bent. Squamous hair	CS
	light brown. Larger, 13–14 mm	
_	Tarsal segment 5 of the hind leg shorter, at most as long as the two preced	
	ing segments taken together, only weakly bent. Squamous hairs darker gre	•
	brown. Somewhat smaller, not larger than 12 mm	
8	Squamous hairs very thick, in fresh specimens the underlying scutal punc	
	tures are obscured. Midline of the scutum is only slightly impressed. Process	
	of labrum markedly elongate with a clear apical emargination in the for	
	margin. Larger, 10–12 mm	
_	Squamous hairs moderately thick, the underlying scutal punctures clear	
	ly visible. Midline of the scutum clearly impressed. Process of labrur	
	regularly trapezoidal, not elongate, fore margin almost straight. Smalle	
0	8–10 mm paucisquama Noskiewic	
9	Tibial scopae composed predominantly of simple hairs, at most with occa	
	sional and scattered plumose hairs (Fig. 64G)	
_	Tibial scopae extensively composed of plumose hairs (Fig. 64H)1	3

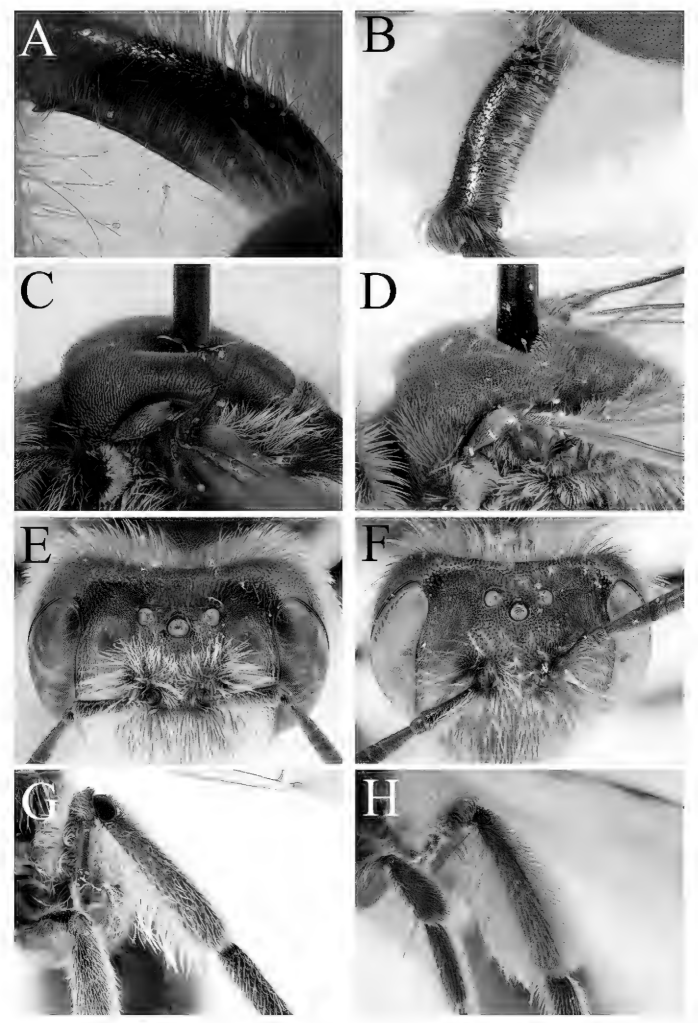


Figure 64. Andrena (Chlorandrena) humilis Imhoff, 1832, female A posterior face of hind femur; Andrena (Rufandrena) orbitalis Morawitz, 1871, female B posterior face of hind femur H tibial scopa, profile view; Andrena (Lepidandrena) pandellei Pérez, 1895, female C scutum, profile view; Andrena (Chlorandrena) leucolippa Pérez, 1895, female D scutum, profile view E foveae, dorsal view; Andrena (Lepidandrena) curvungula Thomson, 1870, female F foveae, dorsal view; Andrena (Cryptandrena) ventricosa Dours, 1873, female G tibial scopa, profile view.

10	Propodeal triangle poorly defined, without lateral carinae, smooth and with
	granular shagreen, more or less shiny over the majority of its area. Larger,
	9–10 mm
_	ally and medially. Smaller, length never exceeding 8 mm
11	Terga strongly and densely punctate, punctures separated by 0.5 puncture
11	diameters, underlying surface smooth and shiny. Propodeal triangle clearly
	delineated laterally by strongly raised carinae ventricosa Dours
_	Terga shagreened and matt, with obscure and shallow punctures. Propodeal
	triangle lacking lateral raised carinae, poorly defined
12	Process of the labrum narrow, clearly produced into a narrow, apically
12	pointed triangle. Head and mesosoma with black pubescence
_	Process of the labrum broad, clearly at least three times broader than long.
	Head and mesosoma with brownish to greyish-white pubescence
	avara Warncke aggregate (potentially including multiple valid species)
13	Scutum and scutellum with clear longitudinal striationsrhyssonota Pérez
_	Scutum and scutellum without longitudinal striations14
14	Fovea with inner margin clearly dorsally diverging from the inner margin of
	the compound eye, curved towards the lateral ocellus (livens-group)15
_	Fovea with inner margin not dorsally diverging from the inner margin of the
	compound eye, not noticeably curved towards the lateral ocellus17
15	Terga with weak but clear metallic green-blue reflections. Antennae usually
	ventrally dark, at most obscurely lightened dark brown. Process of the labrum
	trapezoidal, small, only lightly broader than longnigroolivacea Dours
_	Terga dark, without any metallic reflections. Antennae usually extensively
	lightened orange ventrally. Process of the labrum broader, at least two times
1.	wider than long
16	Discs of T2-3 at their bases consistently and densely punctate, punctures
	typically separated by 1 puncture diameter, at most by 2 puncture diameter.
	eters. Distributed throughout Iberia
_	Discs of T2–3 at their bases more sparsely punctate, punctures separated by
	more than 3 puncture diameters. Rare, known only from a single specimen
17	captured near Madrid
1/	gellum (<i>taraxaci</i> -group)18
	Foveae not strongly constricted ventrally, more or less as wide as dorsally,
	not narrower than the width of a flagellum20
18	Scutum medially with greatly reduced shagreenation, here more or less
10	smooth and shiny. Facial pubescence relatively dark, with black to dark
	brown hairs along the inner margins of the compound eyes. Rare, southern
	and south-eastern Spain only
_	Scutum shagreened, uniformly dull across its entire surface. Facial pubes-
	cence bright, with at most scattered dark hairs. Throughout Iberia 19

19	Depressions of terga, especially T3–4, extensively lightened orange-yellow, semi-transparent. Discs of T2–3 densely punctate, punctures with strongly raised rims
_	Depressions of terga at most with apical rim narrowly lightened orange-yellow. Discs of terga weakly and relatively obscurely punctate, puncture rims weakly raised
20	Tergal margins entirely lightened whitish-yellow hyaline. Face (particularly vertex), scutum, and scutellum with subtle metallic blue-green reflections21
_	Tergal margins never entirely lightened, at most narrowly lightened yellow hyaline. Face (particularly vertex), scutum, and scutellum dark, without metallic reflections
21	Found in dry to steppic areas in central Spain. Scutum with clear punctures that are visible against the strong background shagreenation. Scutellum with shiny interspaces between punctures ³ elata Warncke
_	Found in areas close to or on the coast in southern Portugal and Spain. Scutum with obscure punctures that disappear into the strong background shagreenation. Scutellum with interspaces shagreened ³
22	T2–4 depressed at their base, marginal areas of T2–4 with apical white hair bands that obscure the underlying surface in fresh specimens. Posterior face of the hind femur with long spines
_	T2–4 not depressed basally, apically with marginal areas with at most scattered orange-yellow hairs, these not forming distinct bands which obscure the underlying surface. Posterior face of the hind femur with short teeth
23	T2 between the punctures weakly shagreened, comparatively shiny. Larger species, over 10 mm. Found in more temperate parts of Iberia, generally absent in hot Mediterranean habitats ⁴
_	T2 between the punctures strongly shagreened and dull. Smaller species, under 10 mm. Found in hotter parts of Iberia, the dominant taxon in Mediterranean habitats ⁴
24 (1)	Hind tibial spur clearly broadened, either at its base (Fig. 65A) or submedially (Fig. 65B). Large to very large species, at least 12 mm in length
_	Hind tibial spur not broadened at its base or submedially, more or less parallel-sided. Size variable
25	Propodeum dorsolaterally with dense punctures, punctures separated by <1 puncture diameter (Fig. 65C). Propodeum rounded in profile view, without clearly separated horizontal and vertical parts (subgenus <i>Holandrena</i> Pérez)26
_	Propodeum dorsolaterally with raised reticulation or rugosity (Fig. 65D), without dense punctures or without punctures at all. Propodeum with separated horizontal and vertical parts

³ These two species are challenging to separate in the female sex; geographical context cues and association with the more distinctive males should be used.

⁴ These two species are challenging to separate in the female sex. Association with the more distinctive males should be made.

26	Vertex wide, ocelloccipital distance at least as wide as three times the diameter of a lateral ocellus
_	Vertex narrower, ocelloccipital distance never as wide as three times the di-
	ameter of a lateral ocellus
27	Disc of T1 densely punctate, punctures typically separated by 1 puncture
	diameter. Scutellum dull between punctures. Tergal hair bands interrupted
	medially
_	Disc of T1 more sparsely punctate, punctures separated by greater than 1
	puncture diameter, typically by 2 puncture diameters. Scutellum shiny be-
	tween punctures. Tergal hair bands complete in fresh specimens28
28	Flying in the spring (April–May). Terminal fringe dark blackish-brown.
	Ocelloccipital distance narrower, 1 times the diameter of a lateral ocellus ⁵
	flavilabris Schenck
_	Flying in the summer (July–August). Terminal fringe light. Ocelloccipital dis-
20	tance wider, 1–1.5 times the diameter of a lateral ocellus ⁵ <i>decipiens</i> Schenck
29	Hind tibial spur broadened submedially. Propodeal triangle simply defined
	laterally by weak and obscurely raised carinae, internal surface without a dense network of strongly raised honeycomb-like rugosity. Terga often red-
	marked. Exclusively summer flying species, from May onwards30
_	Hind tibial spur broadened at its base. Propodeal triangle clearly defined
	laterally by strongly raised carinae (Fig. 65D), internal surface with dense
	network of strongly raised honeycomb-like rugosity (though this can be
	weak in members of the subgenus Suandrena Warncke). Tergal only very
	rarely red-marked. Flying in the spring or summer31
30	Tibial scopa ventrally composed of long plumose hairs. Posterior face of the
	hind femora with clear latitudinal carina. Terga usually red marked, though
	an entirely melanic form can be found in south-eastern Spain (ssp. nigricauda
	Wood). Associated with scabious (former Dipsacaceae = Caprifoliaceae). Re-
	stricted to montane grasslands in northern and central Spain with isolated pop-
	ulations in the Sierra de Cazorla and Sierra Nevada hattorfiana (Fabricius)
_	Tibial scopa simple or with at most occasional obscurely plumose hairs.
	Posterior face of the hind femora without a latitudinal carina. Terga never
	red-marked. Associated with yellow Cichorioideae (Asteraceae). Restricted
21	to the Pyrenees and Cantabrian Mountains
31	Terminal fringe dark medially and white laterally (Fig. 24D, F). Lateral faces
	of the propodeum with strongly pronounced network of coarse ridges and
	Terminal fringe uniformly dark. Lateral faces of the propodeum without
_	ridges, either unsculptured or at most with individually raised points which
	do not joint together to form a network34
	do not joint together to form a network

⁵ These two species can be challenging to separate. Association with males should be made, along with phenological context.

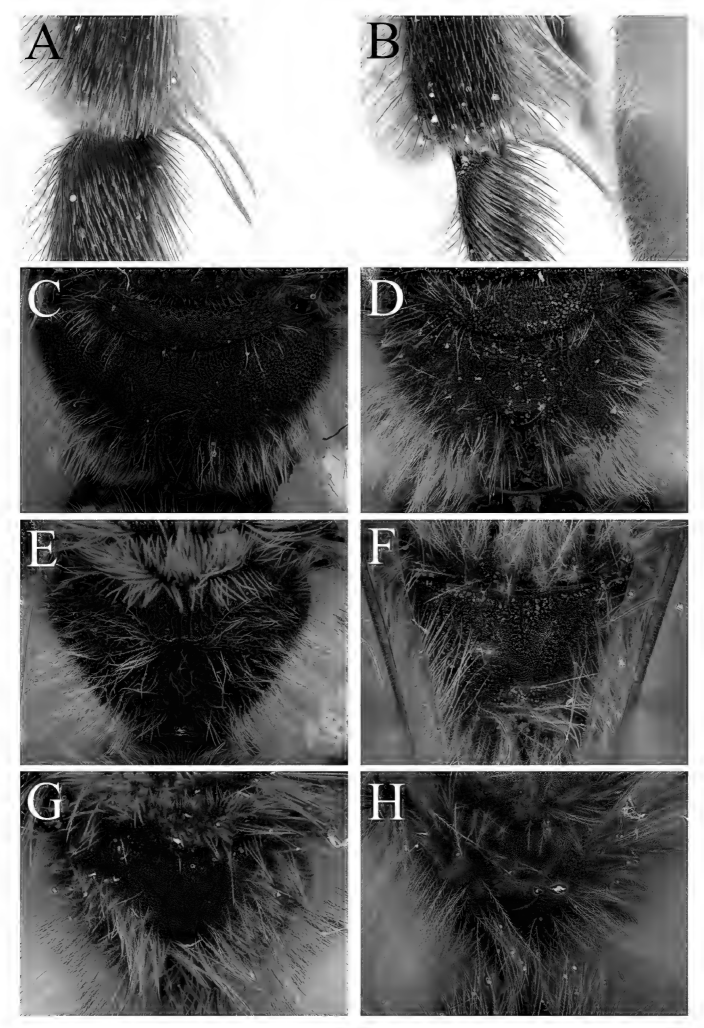


Figure 65. Andrena (Plastandrena) bimaculata (Kirby, 1802), female **A** hind tibial spurs, dorsal view **D** propodeum, dorsal view; Andrena (Holandrena) labialis (Kirby, 1802), female **B** hind tibial spurs, dorsal view **C** propodeum, dorsal view; Andrena (Trachandrena) haemorrhoa (Fabricius, 1781), female **E** propodeum, dorsal view; Andrena (Melandrena) nigroaenea (Kirby, 1802), female **F** propodeum, dorsal view; Andrena (Andrena) helvola (Linnaeus, 1758), female **G** propodeum, dorsal view; Andrena (Hoplandrena) scotica Perkins, 1916, female **H** propodeum, dorsal view.

32	Disc of T1 strongly and densely punctate, punctures separated by <1 punc-
	ture diameter
_	Disc of T1 with sparse punctate, punctures on average separated by 2 puncture diameters
33	Metasoma with metallic blue reflections. T4 laterally with thick white hair
33	patches. Pygidial plate narrow, laterally with a shiny depressed marginal
	area
_	Metasoma black, without metallic blue reflections. T4 laterally with only
	loose white hair, not forming dense hair patches. Pygidial plate broad, flat, without a depressed and shiny marginal area
	afrensis Warncke
34	Terga with metallic blue-green reflections. Mesepisternum and dorsolateral
J 4	
	faces of the propodeum more or less smooth, granularly shagreened, weakly
	and obscurely punctate (subgenus <i>Suandrena</i>)
_	Terga black or red-marked, but never with metallic reflections. Mesepister-
	num and dorsolateral faces of the propodeum with dense network of raised
	rugosity37
35	Propodeal triangle weakly defined, internal rugosity fine and obscure.
	Known only from Cádiz province, flying in December
	gades Wood & Ortiz-Sánchez
_	Propodeal triangle strongly defined, with pronounced internal rugosity.
	More widespread across Iberia
36	Process of the labrum triangular with clear apical point. Throughout Ibe-
	ria ⁶ suerinensis Friese
_	Process of the labrum broadly triangular with apical margin truncate. Con-
	firmed males known only from central, eastern, and south-eastern Spain ⁶
27	
37	Pronotum laterally with strong humeral angle. Terga with extremely fine
	punctation, punctures minute, separated by <0.5 puncture diameters
_	Pronotum laterally rounded, without humeral angle. Terga with much
	coarser punctation, punctures separated by >1 puncture diameter38
38	Pubescence of body predominantly black, tibial scopa predominantly com-
	posed of white hairs39
_	Pubescence of body variable, but usually with extensive brown hairs on mes-
	osoma; tibial scopa usually orange, never predominantly composed of white
	hairs40

⁶ These two species cannot be adequately separated in the female sex. Warncke's unpublished key gave this character, but I am not convinced that it is robust. Association must be made with the distinctive males.

39	Bivoltine (typically March–April and July–August). Common and widely distributed throughout Iberia ⁷
_	Univoltine (typically May–June). Rare, restricted to mountainous parts of Iberia; known from the Pyrenees, the Sistema Central, Serra do Gerês,
40	Sistema Ibérico, and Sierra de Cazorla ⁷
_	Depression of T1 densely punctate, punctures separated by 0.5 puncture diameters. Bivoltine, flying in the spring and the summer ⁸
4 - 4 - 4	
41 (24)	Hairs of the tibial scopae clearly plumose, with a majority of strongly branched hairs
_	Hairs of tibial scopae entirely simple, or with at most a mixture of simple and scattered and weakly branched hairs
42	Terga metallic blue, densely punctate, punctures separated by 1 puncture
_	diameter
42	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
43	Terga with discs extensively red-marked (some dark individual of <i>A. marginata</i> Fabricius with red markings restricted to the base of the tergal margins)
_	Terga dark, at most with depressions lightened, never with red markings at the base of the tergal depressions
44	Clypeus with fore margin slightly upturned. Foveae short, ventrally reaching only to the level of the antennal insertions. Tibial scopae uniformly yellow-
	ish
_	Clypeus with fore margin straight, not upturned. Foveae long, ventrally extending beyond the level of the antennal insertions. Tibial scopae bicol-
	oured, black-brown dorsally, brownish-white ventrally. Associated with sca-
	bious (former Dipsacaceae = Caprifoliaceae) marginata Fabricius
45	Tibial scopae with short hairs, at most half the length of the diameter of the hind tibia at its maximum apical width ranunculi Schmiedeknecht
_	Tibial scopae with long hairs, clearly exceeding half the length of the diam-
46	eter of the hind tibia at its maximal apical width

⁷ These two taxa cannot be separated morphologically. Association must be made with males, or genetic barcodes generated.

⁸ These two taxa are challenging to separate in the female sex due to extreme variation in colouration. Further genetic work is needed to clarify species boundaries in Iberia and more broadly. Association should be made with males.

	Mid and hind basitarsi and hind tibiae lightened orange-yellow (subgenus <i>Chrysandrena</i> Hedicke)
_	Fovea broader, dorsally occupying at least ½ the space between the compound eye and the lateral ocellus, not narrowed ventrally. Basitarsi and hind tibiae dark
47	Clypeus and scutellum strongly shagreened and matthesperia Smith
_	Clypeus and scutellum at least partially shiny
48	Disc of scutum shiny. Scutellum with uniformly light hairs. A2 as long as A3+4. Restricted to temperate areas in northern Portugal and Spain
_	Disc of scutum shagreened and dull. Scutellum with intermixed light and dark brown to black hairs. A2 as long as A3+4+5. Found in Mediterranean areas from southern Portugal to southern and eastern Spain
, -	
49	Process of the labrum large, as long as broad, ventral surface covered with latitudinal wrinkles (Fig. 73D). Foveae dorsally poorly defined, not strongly impressed. Terminal fringe composed of simple dark hairs. Flying in the summer (June-August), associated with Dipsacaceae (=Caprifoliaceae)
_	Process of the labrum twice as broad as long, without wrinkles. Foveae dorsally strongly impressed and therefore well-defined. Terminal fringe composed of densely plumose orange hairs. Flying in the spring (April-May), Associated with yellow Cichorioideae (Asteraceae)
50 (41)	Propodeal triangle clearly defined by strongly raised carinae, internal surface rugose-areolate (Fig. 65E, c.f. Fig. 79A). Mesepisternum and dorsolateral parts of the propodeum with dense network of raised rugosity
_	Propodeal triangle not strongly defined by lateral carinae with its internal surface rugose-areolate (Fig. 65F–H). Mesepisternum and dorsolateral parts of the propodeum with or without dense network of raised rugosity
51	Forewing with two submarginal cells. Clypeus with longitudinal striations.
_	Forewing with three submarginal cells. Clypeus without longitudinal striations
52	Hind tibiae usually orange. Foveae clearly medially constricted. Terga without hair bands, T2 laterally without a pair of small foveae. Terminal fringe
	orange
_	Hind tibiae always dark. Foveae lacking medial constriction. Terga in fresh specimens with apical white hair bands, T2 laterally with a pair of small foveae. Terminal fringe whitish-yellow
53	Process of the labrum clearly deeply emarginatecolletiformis Morawitz
_	Process of the labrum triangular, anterior margin truncate, never emargin-
	ate

54 (50)	Scutum and scutellum with short squamous hairs (Figs 33C, D, 43A, 45C,
	46B)
_	Scutum and scutellum without squamous hairs. If specimen shows longer
55	semi-squamous hairs (e.g. A. farinosa Pérez, Fig. 45D), continue here63
55	Ocelloccipital distance at least three times the diameter of a lateral ocellus
	(Figs 33B, 43B). Large bees, exceeding 11 mm in length
_	Ocelloccipital distance less than two times the diameter of a lateral ocellus.
	Smaller bees, exceptionally reaching 11 mm in length60
56	Dorsolateral faces of the propodeum clearly, densely, and deeply punctate,
	punctures separated by <0.5 puncture diameters, with shiny interspaces57
_	Dorsolateral faces of the propodeum either impunctate, with dense network
	of raised rugosity, or with shallow and sparser punctures, punctures sepa-
	rated by 0.5–2 puncture diameters, interspaces dull
57	Pubescence of scutum weakly squamous, anterior dorsolateral corners of scu-
	tum with pubescence longer, clearly exceeding width of antennae in length.
	Terga always predominantly red. Terga with clear apical hair bands. Puncta-
	tion of T1 slightly spaced, punctures separated by 1-2 puncture diameters.
	Restricted to eastern and south-eastern Spainnilotica Warncke
_	Pubescence of scutum short and strongly squamous, anterior dorsolateral
	corners of scutum with pubescence short, clearly shorter than width of an-
	tennae. Terga variable, red to black or any intermediate combination. Terga
	with or without clear apical hair bands. Punctation of T1 denser, punctures
	separated at most by 1 puncture diameter58
58	A3 exceeding A4+5 in length. Terga always extensively red-marked (central
	and south-eastern Spain)parata Warncke
_	A3 equalling A4+5 in length. Terga variable, from almost entirely black
	(Fig. 33D) to rarely entirely red-marked pruinosa Erichson
59	Scutal hairs orange-brown (Fig. 43A). Hind tibiae orange (Fig. 43C). Terga
	with obscure and narrow brownish-yellow hair bands (Fig. 43D). Terminal
	fringe orange
_	Scutal hairs black and whitish-brown. Hind tibiae dark. Terga with clear
	and thick white apical hair bands. Terminal fringe dark brown
60	T1 strongly and densely punctate, interspaces shiny (Fig. 46D). Mesepister-
	num evenly rounded ventrallyoviventris Pérez
_	T1 weakly and obscurely punctate, underlying surface shagreened.
	Mesepisternum ventrally with an indentation anterior to the attachment
	point of the mid leg61
61	Clypeus without latitudinal ridges. T3–4 laterally with clear squamous hairs
	between the disc and marginal areas. Terga usually extensively red-marked.
	Basitarsi and hind tibiae orange
_	Clypeus with latitudinal ridges. T3–4 laterally without squamous hairs.
	Terga never red marked. Legs dark. 62

62	T2–3 laterally strongly shagreened, with obscure and scattered punctures
_	T2–3 laterally finely shagreened, clearly and densely punctate, punctures separated by 0.5 puncture diameters
63 (54)	Viewed laterally, propodeal corbicula with internal surface (lateral faces of the
00 () -)	propodeum) glabrous AND propodeal corbiculae complete (possessing both
	a dorsal and anterior fringe) (Fig. 66A; subgenus <i>Simandrena</i>)
	Propodeal corbiculae either incomplete (possessing only a dorsal fringe)
_	
(/:	AND/OR internal surface with hairs
64	Tibial scopae bicoloured, dorsally dark and ventrally pale
_	Tibial scopae unicolourous, dorsally and ventrally light
65	Fovea extremely broad, occupying entirety of space between the compound
	eye and the lateral ocellus (Fig. 66B). T1 extremely densely punctate,
	punctures separated by <0.5 puncture diameters, with shiny interspaces.
	Very rare, known only from one specimen Cádiz rhypara Pérez
_	Fovea narrower, not occupying entirety of space between the compound eye
	and the lateral ocellus. T1 strongly shagreened, impunctate or with scattered
	punctures66
66	T2 impunctate. A2 longer than A3+4, almost as long as A3+4+5
	vetula Lepeletier
_	T2 with fine and dense punctures, punctures separated by 0.5 puncture
	diameters. A2 as long as A3+4
67	Terga with thick white apical hair bands in fresh specimens (Fig. 27D).
	Foveae with outer margin straight, not deviating from the inner margins of
	the compound eyes
_	Terga with at most obscure narrow brownish hair bands. Foveae with outer
	margin constricted, clearly deviating from inner margins of the compound
	eyes submedially (Fig. 66C)
68	Tibial scopae with short hairs, dorsally these hairs not greatly exceeding the
00	width of a lateral ocellus. Metasoma with punctures on tergal discs dense
	medially, becoming sparse laterally69
	Tibial scopae with long hairs, very clearly greatly exceeding the width of a
_	
	lateral ocellus. Metasoma with consistently dense punctures, not becoming
(0	sparser laterally
69	Scutum medially strongly shagreened and dull (Fig. 66D). In fresh speci-
	mens, tergal hair bands short, not noticeably surpassing the apex of the
	tergal margins. Hind tibiae and basitarsi usually orange, but can be entirely
	dark. Restricted to temperate areas in northern Spaindorsata (Kirby)
_	Scutum medially polished and shiny (Fig. 66E) or extensively but weakly
	shagreened with a silky shine. In fresh specimens, tergal hair bands long,
	clearly surpassing the apex of the tergal margins. Hind tibiae and basitarsi
	usually dark, occasionally lightened orange-brown. Common throughout
	Iberiapropinqua Schenck

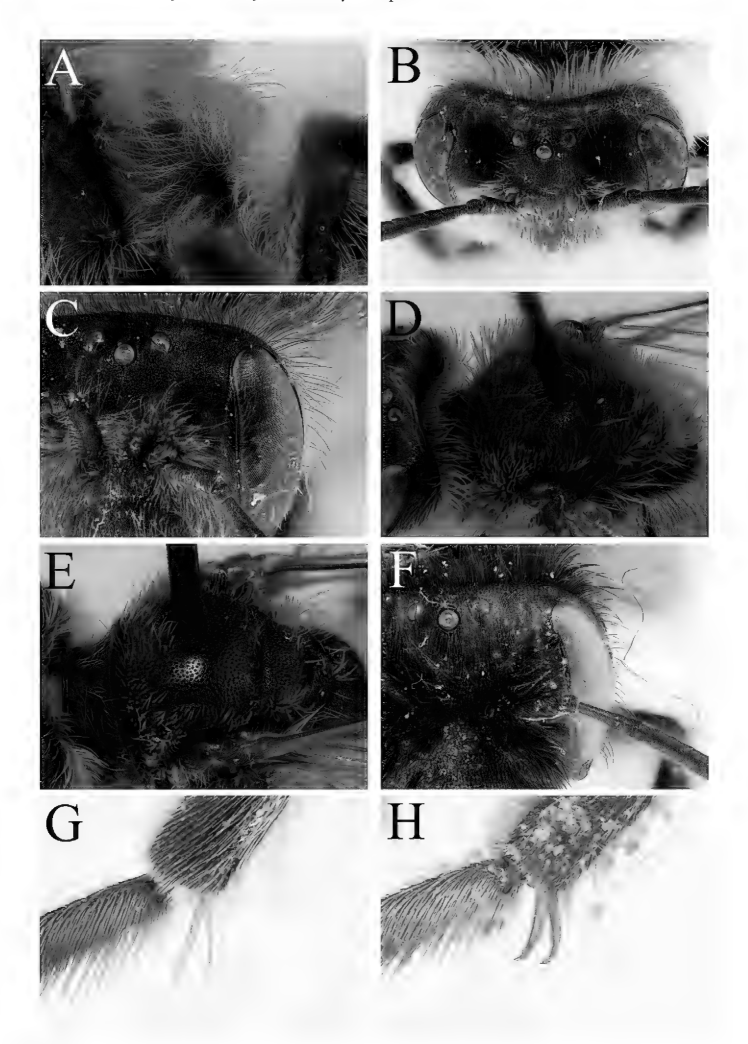


Figure 66. Andrena (Simandrena) dorsata (Kirby, 1802), female **A** propodeal corbicula, lateral view **D** scutum, dorsolateral view; Andrena (Simandrena) rhypara Pérez, 1903, female **B** foveae, dorsal view; Andrena (Simandrena) antigana Pérez, 1895, female **C** fovea, dorsofrontal view; Andrena (Simandrena) propinqua Schenck, 1853, female **E** scutum, dorsolateral view; Andrena (Melandrena) bicolorata (Rossi, 1790), female **F** fovea, dorsofrontal view; Andrena (Notandrena) nigroviridula Dours, 1873, female **G** hind tibial spurs, dorsal view; Andrena (Notandrena) varuga Warncke, 1975, female **H** hind tibial spurs, dorsal view.

70	Disc of T1 strongly and densely punctate, punctures separated by <1 puncture diameter
_	Disc of T1 finely and sparsely punctate, punctures separated by >1 puncture
	diameter
71	Clypeus with pattern of raised latitudinal ridgescombinata (Christ)
_	Clypeus smooth in the middle, at most transversely wrinkled at the base, ventro-laterally with slight longitudinal wrinkles
72	Head and mesosoma white-haired. Clypeus smooth and shiny between punctures. Restricted to areas close to the Pyrenees thomsonii Ducke
_	Head and mesosoma with rich chestnut-brown hair. Clypeus shagreened
73	and dull between punctures
_	T2–4 with discs sparsely and obscurely punctate. Terminal fringe reddish-
	brown. Restricted to the Pyrenees and Cantabrian Mountains ⁹
74 (63)	Fovea strongly constricted medially, strongly diverging from the inner mar-
	gin of the compound eye dorsally (Fig. 66F; former subgenus <i>Hyperandrena</i> Pittioni)
_	Fovea not constricted medially, not strongly diverging from the inner mar-
	gin of the compound eye76
75	Tibial scopae unicolourous black. Mesosoma entirely covered with light grey hairs
_	Tibial scopae dorsally black, ventrally orange-red. Mesosoma entirely cov-
	ered with dark to light brown hairs
76 (74)	At least some tergal discs extensively red-marked (Fig. 41D)77
_	Tergal discs never red-marked, at most with tergal margins lightened
77	Small bees, never exceeding 9 mm in length. Fovea narrow, at most occupy-
	ing ½ of space between the compound eye and the lateral ocellus
_	Large bees, greater than 11 mm in length. Fovea broader, occupying at least
70	½ the space between the compound eye and the lateral ocellus80
78	Propodeal triangle broad, laterally delineated with raised straight carinae, internal surface with fine network of raised rugae. Head and scutum with-
	out metallic reflections
_	Propodeum narrow, poorly defined, lacking lateral carinae, internal sur-
	face at most with short and weak rugae basally, surface with fine granular
	shagreenation. Head and scutum with strong or weak metallic reflec-
	tions

⁹ These two species are challenging to separate, and there may be taxonomic complexity in Iberia. Further study is required.

79	Clypeus medially smooth and shiny between the punctures. Scutum anteriorly shagreened, medially becoming smooth and shiny, irregularly but clearly punctate, with strong greenish metallic reflections
_	Clypeus medially completely shagreened and dull. Scutum shagreened and shallowly punctate, with weak bronzy metallic reflections
80	Tibial scopae unicolourous light orange. Propodeal triangle clearly deline-
80	ated laterally by fine carinae, internal surface with fine network of raised
	rugae. Clypeus densely and uniformly punctate, without a longitudinal im-
	punctate midline
	Tibial scopae bicoloured, dorsally dark, ventrally light (Fig. 41A). Propodeal
_	triangle poorly defined, lacking lateral carinae, internal surface more or less
	smooth, without network of rugae (Fig. 41C). Clypeus densely or sparsely
	punctate, with clear longitudinal impunctate midline81
81	Terga strongly and densely punctate, punctures separated by 1 punc-
01	ture diameter. Compound eyes with inner margins diverging ventrally
	(Fig. 41B). Excluding impunctate midline, clypeus comparatively sparsely
	punctate, punctures separated by 1–2 puncture diameters
_	Terga shallowly and obscurely punctate, punctures separated by 2–3 punc-
	ture diameters. Compound eyes with internal margins more or less paral-
	lel. Excluding impunctate midline, clypeus comparatively densely punctate,
	punctures separated by 0.5–1 puncture diameters 82
82	T1–2 with long hair, equalling the length of the hair on the mesosoma 10
	trimmerana (Kirby) (partim, light form)
_	T1–2 with shorter hair, never equalling the length of the mesosomal hair 10
83 (76)	Head, mesosoma, or tergal discs with metallic reflections (note, A. nigroaenea
	(Kirby) can have bronzy reflections on the terga; if the tibial scopae is com-
	posed of orange-red hairs or the tibial scopae is mainly dark with only the
	ventral hairs orange, go to 206)
_	Body without metallic reflections91
84	Mesosoma with strongly contrasting pattern of black and white hairs; meso-
	soma anteriorly and posteriorly with white hairs, medially with a band of
	black hairs85
_	Mesosoma without strongly contrasting pattern of black and white hairs86
85	Wings infuscate over their apical 2/3rds. Sterna laterally and apically with
	black hairs. A3 comparatively shorter, shorter than A4+5. Bivoltine (April-

¹⁰ Beware of abraded specimens when dealing with these two species.

	May; July-August), restricted to mountains in northern Spain and the
	Pyrenees ¹¹ barbareae Panzer
_	Wings hyaline over the majority of their area, only slightly darkened api-
	cally. Sterna laterally and apically with white hairs. A3 comparatively longer,
	equalling A4+5. Univoltine (April-May), more widespread across northern
	Portugal and Spain ¹¹ cineraria (Linnaeus)
86	Scutum uniformly densely shagreened and completely dull over its entire
	surface, extremely densely punctate, punctures shallow, flat, and confluent.
	Mesosoma and discs of T1-2 dorsally with long chestnut-brown hair
_	Scutum either at least partly shiny or less densely punctate, punctures clearly
	separated by at least 1 puncture diameter. Mesosoma and discs of T1-2
	never with long chestnut-brown hair, either glabrous, with shorter hairs, or
	with hairs of a different colour87
87	Terga clearly punctate, punctures separated by up to 2 puncture diameters.
	88
_	Terga obscurely and sparsely punctate, punctures separated by 4-5 puncture
	diameters89
88	Fovea broad, occupying 2/3rds of the space between the compound eye and
	a lateral ocellus. Tergal punctation comparatively sparse, punctures sepa-
	rated by 1-2 puncture diameters. Scutum medially becoming smooth and
	shiny between the punctures. Restricted to south-western Spain (Huelva,
	Sevilla)
_	Fovea narrow, occupying less than ½ the space between the compound eye
	and a lateral ocellus. Tergal punctation dense, punctures separated by up to
	1 puncture diameter. Scutum uniformly shagreened and dull. Restricted to
	the Pyrenees
89	Larger species, 11-12 mm. Clypeus shagreened and dull over the majority
	of its surface. Fovea narrow, but uniformly wide along their length, not nar-
	rowing ventrally
_	Smaller species, under 9 mm. Clypeus shiny over the majority of its area.
	Foveae narrow, but also narrowing ventrally to approximately half of their
	dorsal width90
90	Scutum with clear metallic green reflections. Hind tibial spurs straight
	(Fig. 66G)nigroviridula Dours
_	Scutum dark, with at most weak and obscure metallic reflections. Hind
	tibial spurs strongly bent at their apexes (Fig. 66H) varuga Warncke
91 (83)	Small black species without a keel laterally on the pronotum, body length
	under 8 mm, or if up to 10 mm then with lateral faces of the propodeum

¹¹ These two species are challenging to separate morphologically.

	with clear pattern of raised star-shaped wrinkles (all members of the subgenera <i>Aciandrena</i> , <i>Graecandrena</i> , <i>Micrandrena</i>) ¹²
_	Species larger than 9 mm in length or with pronounced keel on the pronotum laterally
92	Propodeal triangle smooth, not defined laterally by raised carinae, with internal surface lacking network of raised rugosity, at most with very short
-	rugae at the base of the propodeal triangle (Fig. 67A)
	(Fig. 67B; note, take care with <i>A. tenuistriata</i> , for which the lateral parts of the propodeal triangle present granular shagreen, Fig. 67C) 106
93	Clypeus with clear raised longitudinal striations (Fig. 67D)94
94	Clypeus without any striations
	long as A4+5+6. Central and eastern Spain only
_	Clypeus evenly arched. Process of the labrum narrower, as long as broad. A3 only slightly exceeding A4+5. More widespread
95	Tibial scopa composed of unicolourous light hairs. T3 impunctate, with weak apical yellowish hair band, marginal area at most weakly depressed. Widespread across Iberia
-	Tibial scopa pale ventrally, dark dorsally. T3 with obscure punctures, with strong apical white hair band overlying the clearly depressed marginal area. Restricted to sandy and usually coastal habitats in southern Iberia
96	Foveae not ventrally narrowed, in their lower half at least half as wide as the distance from the inner margin of the compound eye (Fig. 67E)97
_	Foveae ventrally narrowed, in their lower half as wide as the distance from the inner margin of the compound eye (Fig. 67F, G)
97	T2–4 with narrow, widely interrupted hair bands. Wing venation brownish.
	Clypeus shagreened to smooth and shiny, densely punctate, punctures separated by 1 puncture diameter. Process of the labrum narrow and triangular
	with a pointed tip (Fig. 67E). Larger, 8 mm. Widespread across Iberia
	T2 4 with complex wide and dense white heir hands. Wing wonetien light
_	T2–4 with complex wide and dense white hair bands. Wing venation light yellow. Clypeus uniformly shagreened, irregularly punctate, punctures sepa-
	rated by 1–3 puncture diameters. Process of the labrum with the apical
	margin truncate. Smaller, 6 mm. Rare, known only from central Spain
	montarca Warncke

¹² Note, this couplet is not intended to include small examples of members of the subgenera *Leucandrena* Hedicke, 1933 and *Notandrena*. Therefore, small individuals with a clearly keeled pronotum should follow the alternative couplet.

98	Tergal discs clearly and regularly punctate, punctures separated by 1 punc-
	ture diameter, with punctures extending onto marginal areas
_	Terga either impunctate or obscurely punctate, but punctures never extending onto marginal areas99
99	Supraclypeal area covered with longitudinal striations (Fig. 67F)100
_	Supraclypeal area without any striations (Fig. 67G)
100	Longitudinal striations on the paraocular areas strong and pronounced, contin-
100	uing ventrally to the lateral margins of the clypeus without becoming weaker.
	Clypeus flattened, with distinct longitudinal impression or furrow medially.
	Restricted to the extreme north-east of Spain <i>impunctata</i> Pérez
_	Longitudinal striations on the paraocular areas ventrally extending to the
	lateral margins of the clypeus but here clearly weaker than their strength
	adjacent to the foveae. Clypeus either domed or if flattened then without
	longitudinal impression
101	\mathcal{E} 1
101	Clypeus strongly flattened, coarsely shagreened and dull over almost its
	entirely surface, apical margin narrowly and obscurely shiny; clypeus with
	shallow and obscure punctures that disappear into the underlying sha-
	green, punctures separated by 1–2 puncture diameters (Fig. 67F)
	Claracia side and a series of the control of the co
_	Clypeus either domed, or if flattened then with a broad shiny apical margin
	(at least as broad at the width of a flagellum) and punctures that are clearly
102	visible against the underlying shagreenation
102	A3 exceeding the length of A4+5. Striations of supraclypeal area weakly
	continue onto clypeus basally and laterally, here shagreened with weakly
	raised striations. Clypeus shagreened basally and laterally, becoming smooth
	and shiny medially and apically, with narrow medial shagreened projection,
	shagreenation thus forming a weak trident shape. Known only from a few
	specimens from the extreme south of Spain, probably representing an unde-
	scribed species (Málaga, Sevilla)
_	A3 at most equalling A4+5. Striations of supraclypeal area not continuing
	onto clypeus, entire clypeus free of even a hint of striations103
103	Clypeus with fine punctures, punctures separated by 1–3 puncture diameters.
	Underlying shagreenation weak basally. Process of the labrum slightly broader
	than long. Presence and distribution in Iberia unclear
_	Clypeus with strong and coarse punctures, punctures separated by 0.5–2
	puncture diameters. Underlying shagreenation strong and coarse basally.
	Process of the labrum narrow, slightly longer than broad
104	Scutellum polished and shiny between punctures. Clypeus domed and
	somewhat flattened medially. Restricted to south-eastern Spain (Alicante,
	Almería, Granada, Murcia) tenostra Warncke
_	Scutellum dull and shagreened. Clypeus largely flattened across its entire
	surface. More widespread across Iberiaalma Warncke

105	Larger, 7–8 mm. T3–4 with discs obscurely punctate. Hind basitarsi lightened orange. Nervulus of the forewing interstitial to weakly antefurcal. Sha-
	greenation of the clypeus becoming weaker at its apical margin
_	Smaller, 4–5 mm. T3–4 with discs impunctate. Hind basitarsi dark. Nervulus
	of the forewing strongly antefurcal. Clypeus uniformly shagreened
	vacella Warncke
106	Lateral shoulders of T1 with a pair of strongly produced sharp ridges
100	(Fig. 67H). Terga shiny with mixture of small and large punctures. Mesepis-
	ternum depressed above the insertion point of the mid legs. Rare, restricted
	to central Spain
	Lateral shoulders of T1 never with clearly produced ridges, or if with ridges
_	(A. strohmella Stöckhert) then terga never shiny and deeply punctate. Tergal
	punctation otherwise. Mesepisternum evenly rounded107
107	Hind tibiae and basitarsi orange, tibial scopa composed of extremely short
10/	hairs. Restricted to the Pyrenees and the Cantabrian Mountains, flying
	July–August, associated with <i>Potentilla</i> (Rosaceae) <i>tarsata</i> Nylander
_	Hind legs dark, tibial scopa with hairs normal, not extremely short 108
108	Foveae long, dorsally extent reaching a line parallel to the hind margin of the
100	lateral ocelli, foveae deeply impressed. Propodeal triangle not laterally deline-
	ated by carinae, internal surface weakly elevated, with irregular raised rugae
	that do not cover the entire area. Terga laterally with loose, white interrupted
	hair bands. Restricted to cooler areas in and around the Pyrenees and Canta-
	brian Mountains, flying July–August
_	Foveae shorter, not reaching level of the lateral ocelli dorsally, only weakly
	impressed. Propodeal triangle clearly delineated laterally by carinae, internal
	surface evenly and regularly covered by fine network of raised rugae. Terga
	with lateral hair bands or not
109	Foveae either uniformly narrow or strongly narrowed ventrally, in their ven-
	tral half at most as wide as the distance to the inner margin of the com-
	pound eye110
_	Foveae not or only weakly narrowed ventrally, in their ventral half clearly
	wider than the distance to the inner margin of the compound eye115
110	Clypeus with longitudinal striations (c.f. Fig. 67D)111
_	Clypeus without striations112
111	Disc of T1 shagreened and dull. T2-4 with hair bands long, longer than the
	diameter of a flagellum. Scutum uniformly shagreened and dull. Restricted
	to the extreme north-east of Spain
_	Disc of T1 polished and shiny, contrasting the shagreened marginal area.
	T2-4 with short hair bands, shorter than the diameter of a flagellum. Scu-
	tum laterally shagreened, medially smooth and shiny. Widespread through-
	out Iberianitidula Pérez

112	T1 finely shagreened to polished and shiny. Mesonotum densely and strong-
	ly punctate
_	T1 strongly shagreened and dull. Mesonotum finely and sparsely punctate.
113	Foveae uniformly narrow, dorsally narrower than the diameter of a flagel-
	lum. Disc of T1 with scattered punctures, separated by at least 2 punctures
	diameters
_	Foveae dorsally broader than ventrally, dorsally as wide as the diameter of a
	flagellum. Disc of T1 more densely punctate, punctures separated by 0.5–1
	puncture diameters
114	Foveae uniformly narrow, dorsally narrower than the diameter of a flagel-
	lum. Terga impunctate. Clypeus evenly domed, densely shagreened and
	dull. Associated with Cistaceae
_	Foveae dorsally broader than ventrally, dorsally as wide as the diameter of
	a flagellum. Terga obscurely punctate. Clypeus weakly three-faced, anterior
	margin shiny. Associated with Brassicaceae tenuistriata Pérez
115	Large species, 10 mm in length. Clypeus densely covered with latitudinal
	wrinkles. Lateral faces of the propodeum covered with pattern of raised star-
	shaped wrinkles
_	Most species smaller. Clypeus without latitudinal wrinkles and propodeum
	without pattern of raised star-shaped wrinkles116
116	Larger species, 8-9 mm. Discs of T1-3 extremely densely punctates
	punctures almost confluent, separated by <0.5 puncture diameters. Cl-
	ypeus densely punctate, punctures separated by 1 puncture diameter
	interspaces with weakly raised ridges that form subtle pattern of longi-
	tudinal striations. Rare, known from eastern Spain (Jáen, Soria, Teruel)
	vaulogeri Pérez
_	Smaller, usually shorter than 8 mm. Discs of T1-3 less densely punctate
	punctures separated by >1 puncture diameter, or if punctures dense, then
	much smaller than 8 mm in length. Clypeus without pattern of longitudinal
	ridges or striations117
117	T3-4 with the marginal areas strongly depressed relative to the discs (Fig.
	68A–E)118
_	T3-4 with the marginal areas only superficially depressed (Fig. 68F, G)
118	T2–3 laterally without a gradulus
_	T2–3 laterally with gradulus present (c.f. Fig. 68F, G)
119	Tergal discs and marginal areas completely smooth and shiny, without
	any shagreenation (Fig. 68B). Found in the Sistema Central to west and
	north-western Iberia. Associated with Sedum (Crassulaceae)
	omnilaevis Wood
_	At least tergal discs shagreened

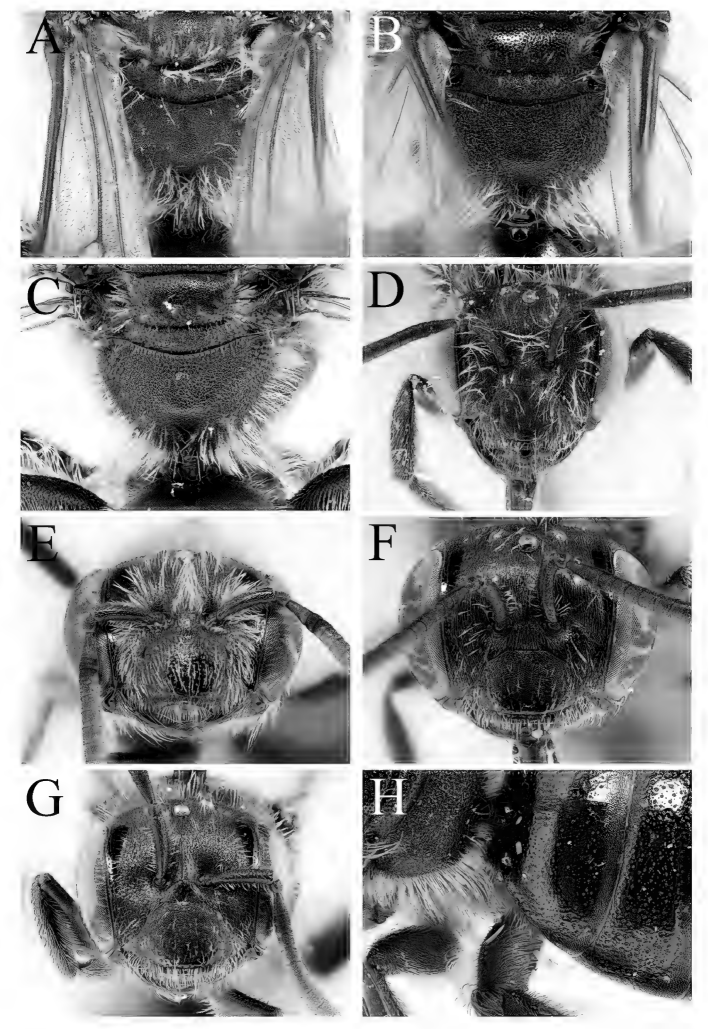


Figure 67. Andrena (Micrandrena) longibarbis Pérez, 1895, female A propodeum, dorsal view D clypeus, frontal view; Andrena (Micrandrena) spreta Pérez, 1895, female B propodeum, dorsal view; Andrena (Micrandrena) tenuistriata Pérez, 1895, female C propodeum, dorsal view; Andrena (Micrandrena) pandosa trigona Warncke, 1975, female E face, frontal view; Andrena (Graecandrena) verticalis Pérez, 1895, female F face, frontal view; Andrena (Graecandrena) nebularia Warncke, 1975, female G face, frontal view; Andrena (Parandrenella) taxana Warncke, 1975, female H T1, dorsal view.

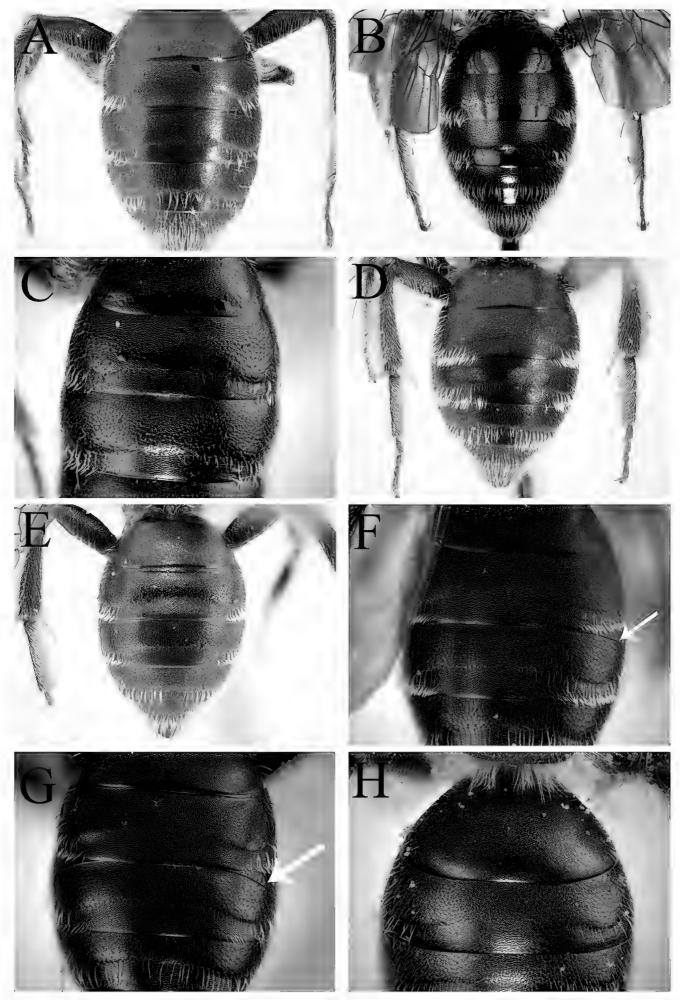


Figure 68. Andrena (Micrandrena) spreta Pérez, 1895, female A terga, dorsal view; Andrena (Micrandrena) omnilaevis Wood, 2020, female B terga, dorsal view; Andrena (Micrandrena) anthrisci Blüthgen, 1925, female C terga, dorsal view; Andrena (Micrandrena) pauxilla Stöckhert, 1935, female D terga, dorsal view; Andrena (Micrandrena) bayona Warncke 1975, female E terga, dorsal view; Andrena (Micrandrena) minutula (Kirby, 1802), female F terga, dorsal view (gradulus indicated by white arrow); Andrena (Micrandrena) minutuloides Perkins, 1914, female G terga, dorsal view (gradulus indicated by white arrow); Andrena (Micrandrena) falsifica Perkins, 1915, female H terga, dorsal view.

120	Terga with marginal areas not noticeably differentiated structurally from discs, strongly and densely shagreened and dull (Fig. 68D). Tergal discs essentially impunctate, with any obscure punctures disappearing into dense shagreenation. Scutum densely punctate, punctures separated by 0.5–1
	puncture diameter
_	Terga with marginal areas noticeably shinier than discs, with weaker sha-
	greenation. Tergal discs with clear punctures visible against the underlying
	shagreenation (Fig. 68C). Scutum less densely punctate, punctures separated by 1–2 puncture diameters
121	Depressed tergal margins polished and shiny, clearly shinier than sha-
121	
	greened tergal discs. In dorsal view, lateral hairs at the base of the marginal
	area of T2 longer, erect, projecting at a 45° angle, breaking profile of meta-
	soma laterally. Margin of T2 comparatively strongly depressed. Restricted
	to the Pyrenees and the Cantabrian Mountains
	semilaevis Pérez
_	Depressed tergal margins shagreened, similar to and not strongly contrast-
	ing shagreened tergal discs (Fig. 68D). In dorsal view, lateral hairs at the base
	of the margin area of T2 shorter, flat, not projecting, almost parallel with
	lateral profile of metasoma. Margin of T2 comparatively weakly depressed.
	Slightly more widespread across central and northern Spain in mountainous
	areasanthrisci Blüthgen
122	Disc of T1 smooth and shiny, clearly punctate, punctures medially separat-
	ed by 0.5-1 puncture diameter, not extending onto the broad impunctate
	marginal area. Rare, restricted to the Pyrenees and northern Spain
	floricola Eversmann
_	Disc of T1 never smooth and shiny, always at least partially shagreened,
	punctate or not
123	Gena entirely smooth and shiny, without any shagreenation. Scutum and
125	scutellum polished and shiny. T2–3 with discs strongly and densely punc-
	tate, punctures separated by 1 puncture diameter (Fig. 68E). Restricted to
	dry and steppic habitats in central Iberia including northern Portugal
_	Gena shagreened and dull, at most with a narrow shiny strip along the
	outer margin of the compound eye. Scutum and scutellum variable, usu-
	ally at least partially shagreened. T2-3 usually less densely and strongly
	punctate
124	Head comparatively long, only slightly shorter than broad (Fig. 48B).
	Discs of T1-3 extremely densely punctate, punctures separated by $0.5-1$
	puncture diameters (Fig. 48D). Restricted to high-altitude on the Sierra
	Nevada ortizi sp. nov.
_	Head shorter, clearly broader than long. T1-3 never so densely punctate,
	punctures when visible separated by at least 1 puncture diameter125

125	Propodeal triangle poorly defined (Fig. 20E, F), lateral carinae disappearing medially, not extending from the base (metanotum) to the apex (vertical face of the propodeum). Tergal discs weakly to strongly shiny
_	Propodeal triangle well-defined, lateral carinae consistently strong from the base to the apex. Tergal discs weakly to strongly shagreened, never shiny
126	Tergal discs entirely shiny, polished, and smooth (Fig. 20H). Scutum and scutellum extensively shiny, without shagreenation
_	Tergal discs at least partially shagreened, never entirely shiny (Fig. 20G). Scutum and scutellum with subtle shagreenation, not smooth and shiny
127	T1 laterally with slightly raised corners, visibly projecting and disrupting rounded profile when viewed dorsally. Clypeus slightly flattened centrally, with clear impunctate mid-line between large and well defined clypeal punctures. Terga 2–3 centrally impunctate, laterally with large punctures with raised margins (crater punctures). Restricted to areas close to the
	Pyreneesstrohmella Stöckhert
_	T1 laterally without such raised projections, in dorsal view evenly rounded. Clypeus with or without a clear impunctate mid-line. Terga punctation with or without lateral crater punctures. Distribution otherwise
128	Clypeus weakly domed, sparsely punctate, punctures large, separated by 2–3 puncture diameters. T2–3 laterally with large hair-bearing punctures that contrast the underlying shagreenation and do not disappear into it <i>icterina</i> Warncke
_	Clypeus more densely punctate, punctures separated by 1–2 puncture diam-
	eters. T2–3 laterally with at most obscure punctures129
129	Terga comparatively weakly shagreened and finely punctate, punctures relatively clearly visible against background shagreen, punctures extending onto tergal margins, most clearly visible on T1–3. Anterior ½ to ⅓ of clypeus usually polished and shiny. Probably univoltine (March-May). Central,
	eastern, and southern Spain, rare ¹³ exigua Erichson
_	Terga comparatively strongly shagreened, obscurely punctate, punctures disappearing into underlying shagreenation (Fig. 68A). Tergal margins impunctate. Clypeus usually shagreened, without polished fore-margin ¹³
130	Scutum strongly and clearly punctate, punctures separated by 1–2 puncture diameters, underlying surface variable but always at least partially shiny. Bivoltine (March-July). Throughout Iberia, common ¹³ spreta Pérez
_	Scutum obscurely punctate, punctures disappearing into the abundant underlying shagreen, scutum dull. Univoltine (April-May). Iberian distribution

¹³ These three species are challenging to separate without confidently identified reference material. Most specimens will be *A. spreta* Pérez which is by far the most commonly collected of the three.

	unclear, but probably restricted to southern Spain (Cádiz, Málaga) ¹³
	tiaretta Warncke
131	Disc of T1 clearly punctate, though sometimes finely and sometimes only
	punctate basally (Fig. 68H). Disc of T2 more strongly, clearly, and exten-
	sively punctate132
_	Disc of T1 finely to strongly shagreened, at most obscurely punctate with
	punctures disappearing into the underlying surface sculpture (Fig. 67F, G).
	Disc of T2 with at most a few punctures laterally
132	Marginal area of T1 wide, strongly thickened, impunctate (Fig. 68H).
	Restricted to mountainous areas in northern Spain, associated with Potentilla
	(Rosaceae) falsifica Perkins
_	Marginal area of T1 normal, not widened or thickened133
133	Disc of T1 strongly shagreened, predominantly punctured medially.
	Antennae ventrally at most lightened dark brown. Throughout Iberia, bi-
	voltinealfkenella Perkins
_	Disc of T1 finely shagreened, evenly and finely punctate over its entire area,
	with shiny interspaces. Antennae ventrally orange. Rare, restricted to north-
	ern Spain, univoltine (August-September)
134	Process of the labrum narrow, as long as broad. Scutum strongly shagreened
-0-	and matt, densely punctate, punctures difficult to discern against the strong
	underlying sculpture. Clypeus sparsely punctate, with obscure and weak
	latitudinal striations
_	Process of the labrum wider, broader than long. Scutum less strongly sha-
	greened, with punctures of variable density that are clearly visible against the
	underlying sculpture. Clypeus variably punctate, but never with latitudinal
	striations
135	Clypeus strongly flattened and sparsely punctate, punctures separated on
137	average by more than 2 puncture diameters. Process of the labrum medially
	emarginate. Associated with <i>Ornithogalum</i> (Asparagaceae)
	Clypeus domed, never strongly flattened, punctures dense or sparse. Process
_	
126	of the labrum truncate or apically rounded
136	Clypeus sparsely and obscurely punctate, punctures separated by 2–3 punc-
	ture diameters or more. Scutum very sparsely and finely punctate, punctures
	small, separated by 2–3 puncture diameters, underlying surface shagreened
	and dull. Restricted to temperate areas close to the Pyrenees
_	Clypeus clearly punctate, punctures separated by 1–2 puncture diameters. Scu-
	tum more densely punctate, punctures larger, separated by up to 2 puncture
	diameters, underlying surface dull to shiny. Throughout Iberia137
137	Scutellum shagreened, at most weakly shiny. Scutum more densely punc-
	tate, punctures separated by 1 puncture diameter, underlying surface sha-
	greened and dull (1st generation) to weakly shiny (2nd generation). Terga
	with dense and narrow apical hair bands laterally, individual hairs touching

	each other in fresh specimens (Fig. 68F). Foveae not noticeably narrowed
	ventrally
_	Scutellum always polished and shiny. Scutum less densely punctate, punc-
	tures less regular, separated by 1-2 puncture diameters, underlying surface
	finely shagreened and weakly shiny (1st generation) to smooth and shiny
	(2 nd generation). Terga with sparse apical hair bands, occasionally with some
	individual hairs touching each other (Fig. 68G; 1st generation), usually with
	all hairs individually separated (2nd generation). Foveae slightly but distinc-
	tively narrowed ventrally
138(91)	Clypeus flattened over majority of its surface. Species strongly associated
	with Fabaceae (subgenus <i>Taeniandrena</i>) ¹⁴ 139
_	Clypeus not noticeably flattened. Species not usually associated with
	Fabaceae
138	Face and foveae black haired. Terminal fringe dark brown (Fig. 69A).
	Terga with weak and obscure punctures. Known only from central Portu-
	gal (Castelo Branco) and south-western Spain (Huelva, Cádiz). Univoltine
	(March–April)
_	Combination of characters otherwise; either face with pale hairs, terminal
	fringe light, or tergal densely punctate140
140	Terminal fringe composed of dark brown to brownish grey hairs. Hairs
	flanking the basitibial plate of the hind tibia also dark (Fig. 69B)141
_	Terminal fringe and hairs flanking basitibial plate of hind tibia light, golden
	to yellow-orange (Fig. 69C–F)143
141	Discs of T2-4 strongly and densely punctate, punctures clearly visible
	against the underling shagreenation. Widespread throughout Iberia, most
	common in areas with an Atlantic climate and abundant Genisteae. Bivolt-
	ine (typically March-May; June-July)ovatula (Kirby)
_	Discs of T2-4 obscurely and shallowly punctate, punctures disappearing
	into background shagreenation. Iberian distribution more restricted (see be-
	low)
142	Scutum with punctures shallow and obscure, separated by >1 puncture di-
	ameter. T3 with apical hair band interrupted medially. Currently known
	only from the steppe of central Spain (Guadalajara, Salamanca, Segovia).
	Univoltine (May–June)
_	Scutum with punctures dense and clear, separated by <1 puncture diam-
	eter. T3 with apical hair band complete in fresh specimens. Currently only
	confirmed from the coast of southern Spain (Málaga). Bivoltine (probably
	February–April; May–June)poupillieri Dours

¹⁴ Females of this subgenus are highly challenging and in some cases impossible to identify. There are several outstanding taxonomic issues in this subgenus that require in-depth genetic study to resolve, and additional probably undescribed cryptic species are present. Do not expect to identify this group without consulting confidently determined reference material, and ideally barcoded specimens. Minimal characters are given here because of the ongoing lack of taxonomic clarity in this subgenus. See also the key of Praz et al. (2022).

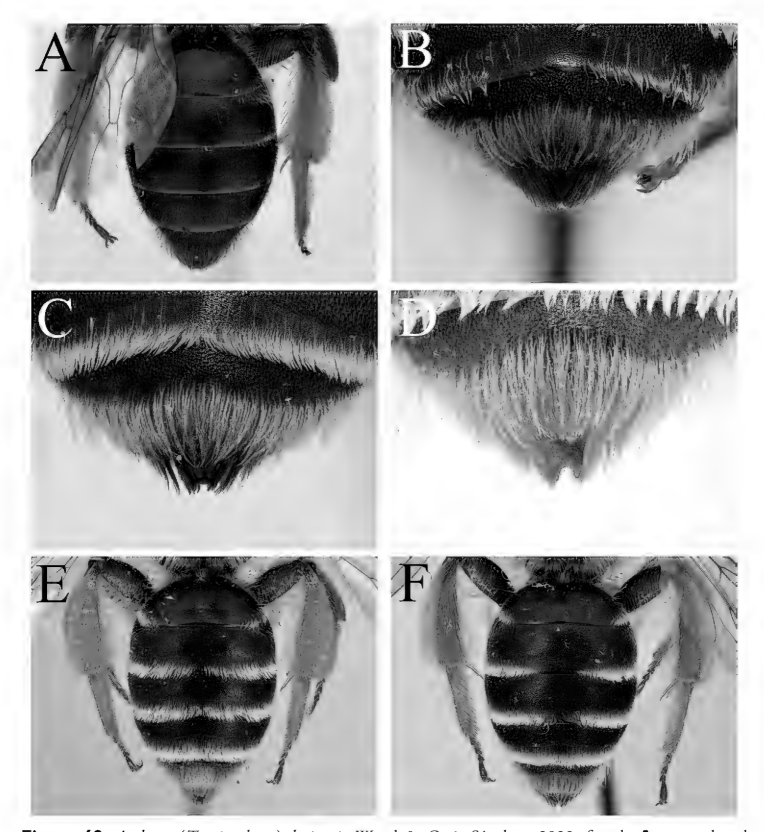


Figure 69. Andrena (Taeniandrena) lusitania Wood & Ortiz-Sánchez, 2022, female A terga, dorsal view; Andrena (Taeniandrena) ovatula (Kirby, 1802), female B terminal fringe; Andrena (Taeniandrena) afzeliella (Kirby, 1802), female C terminal fringe; Andrena (Taeniandrena) lathyri Alfken, 1900, female D pygidial plate, posterior view; Andrena (Taeniandrena) gredana Warncke, 1975, female E terga, dorsal view; Andrena (Taeniandrena) benoisti Wood & Praz, 2021, female F terga, dorsal view.

144	Clypeus apically shiny. T2–4 apically with wide white hair bands that are complete in fresh specimens, these clearly exceeding the length of the marginal areas (Fig. 69E). Scutum laterally shagreened, becoming smooth and
	shiny medially. Restricted to the Sistema Central and northern and western
	Iberia
_	Clypeus uniformly dull. T2–4 with hair bands different, either shorter, broadly interrupted medially, or yellowish (Fig. 69F). Scutum uniformly
	dull or with circular shiny area medially145
145	Declivity of T1 medially strongly and densely punctate, punctures separated by <1 puncture diameter146
_	Declivity of T1 with at most scattered punctures, never with dense patch of
	punctures medially149
146	T3 with interrupted apical hair band in fresh specimens. T2–4 with the apical hyaline part of the marginal areas narrow, not exceeding 3 times the diameter of a puncture from the tergal discs. Scutum uniformly dull. Restricted to northern Portugal and Spain with isolated populations at elevation in the Sistema Central and Sistema Ibérico
_	T3 with apical hair band complete in fresh specimens (Fig. 69F). T2-4 with
	the apical hyaline part of the marginal areas comparatively broad, exceeding 4 times the diameter of a puncture from the tergal discs. Scutum at least
147	partly shiny
14/	and shiny; punctation here becoming weaker and sparser. Restricted to the
	Sistema Central to central and northern Portugal and north-western Spain (Zamora). Not known from the Cantabrian or Pyrenees Mountains
_	Scutum more or less uniformly shagreened and punctate. Found across
	mountainous areas in eastern Spain (Sierra Nevada, Sierra de Cazorla, Sistema Ibérico) to the Pyrenees
148	Restricted to the alpine zone of the Sierra Nevada (above 2000 m)
110	
_	Found elsewhere
	likely represents a complex of an unknown number of species, potentially all
1 40	of which are undescribed; the true <i>intermedia</i> may be absent from Spain)
149	Terga obscurely punctate, at least on the base of T2 with punctures
	disappearing into the underlying shagreenation. Univoltine (April–June)
	russula Lepeletier sensu lato (including the distinct mitochondrial
	lineage from southern Portugal)
_	Terga clearly punctate, at least on the base of T2 with punctures clearly vis-
	ible against the underlying shagreenation150
150	Terga strongly shagreened, dull to weakly shiny. Scutellum shagreened and weakly shiny. Tergal hair bands yellowish. Larger, 11–12 mm. Restricted

	to southern and south-eastern Spain (Almería, Granada, Málaga, Murcia,
	Valencia). Univoltine (April–June)
_	Terga less strongly shagreened, shiny comparatively more strongly. Scutel-
	lum medially almost without sculpture, brightly shiny. Tergal hair bands
	whitish. Smaller, 8-10 mm. Throughout Iberia. Bivoltine (typically May-
	June; July–August)
151 (138)	Fovea dorsally narrow, occupying at most 1/3 of space between lateral ocellus and
, ,	compound eye, ventrally narrowing strongly (subgenus <i>Euandrena</i>) 152
_	Foveae either dorsally broader, or not strongly narrowing162
152	Head elongate, mouthparts extremely long, twice the length of the head
	(c.f. Fig. 89D). Process of the labrum triangular. Associated with Lithodora
	(Boraginaceae)
_	Head and mouthparts shorter, mouthparts never twice the length of the
	head. Process of the labrum trapezoidal
153	Clypeus densely and coarsely punctate with a raised longitudinal impunc-
170	tate shiny line. Restricted to areas close to the Pyrenees. Associated with
	Symphytum (Boraginaceae)symphyti Schmiedeknecht
_	Clypeus densely or weakly punctate, never with a raised longitudinal im-
	punctate shiny line
154	Pronotum with weak but distinct lateral keel. Clypeus medially with shal-
1)1	low longitudinal impression
	Pronotum laterally rounded. Clypeus without longitudinal impression, in
_	, , , , ,
	one species (<i>A. rufula</i> Schmiedeknecht) with hints of an impression caused by impunctate longitudinal midline
155	, 1
155	T2 with marginal area long, occupying almost ½ the segment, strongly de-
	pressed and shiny (Fig. 70A). Facial pubescence light brown, with at most
	dark hairs laterally on the frons. Terminal fringe brown. Found in areas with
	a temperate Atlantic climate across central, north-western, and northern
	Iberia
_	T2 with marginal area shorter, not strongly depressed (Fig. 70B). Face
	with intermixed black hairs throughout, including on the vertex (Fig. 8C).
	Terminal fringe dark brown to black (Fig. 8D). Found in areas with a Medi-
	terranean climate
156	Tergal discs with extremely coarse and dense punctures, punctures sepa-
	rated by 0.5-1 puncture diameter. Tergal margins strongly depressed and
	essentially impunctate, margins of T2-4 overlain by sparse whitish hair
	bands that emerge from the apexes of the tergal discs and which do not ob-
	scure the underlying surface of the marginal areas (Fig. 70C). Known only
	from mountainous areas in north-western, central, and south-eastern Spain
	(Léon, Ávila, Jáen)
_	Tergal discs with normal punctures, not noticeably coarse, punctures separat-
	ed on average by 1 puncture diameter. Tergal margins only weakly depressed,
	without long whitish hair bands, sometimes with obscure brownish lateral
	hair bands on the apexes of the marginal areas themselves157

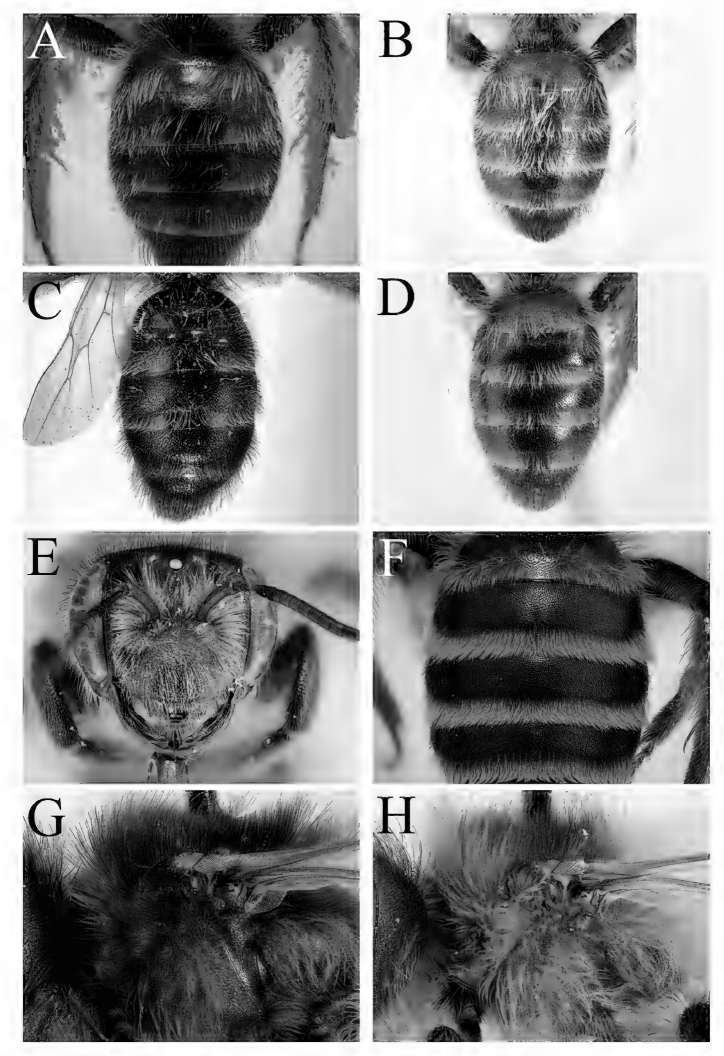


Figure 70. Andrena (Euandrena) angustior (Kirby, 1802), female **A** terga, dorsal view; Andrena (Euandrena) lavandulae Pérez, 1902, female **B** terga, dorsal view; Andrena (Euandrena) fortipunctata Wood, 2021, female **C** terga, dorsal view; Andrena (Euandrena) granulosa Pérez, 1902, female **D** terga, dorsal view; Andrena (Melandrena) flavipes Panzer, 1799, female **E** head, frontal view **F** terga, dorsal view; Andrena (Melandrena) vulcana Dours, 1873, female **G** mesosoma, profile view (dark form) **H** mesosoma, profile view (light form).

157	Apical margins of T1–4 extensively lightened yellow-hyaline, this lightened area clearly exceeding the diameter of a lateral ocellus (Fig. 70D). Associated
_	with Cistaceae
	narrowly so, this lightened area not exceeding the diameter of a lateral ocellus
158	Tergal margins comparatively more strongly depressed. Tergal discs on aver-
	age more coarsely and densely punctate, punctures on the disc of T2 separated by 1.5–2 puncture diameters ¹⁵
_	Tergal margins comparatively more weakly depressed. Tergal discs on aver-
	age less strongly and densely punctate, punctures on the disc of T2 separated
	by 3–4 puncture diameters ¹⁵ <i>vulpecula</i> Kriechbaumer
159	Facial pubescence predominantly light, with dark hairs restricted to the inner
	margins of the compound eyes. Mesepisternum entirely pale-haired. Restricted
	to temperate areas close to the Pyrenees
_	Facial pubescence predominantly black, with occasional scattered light hairs
	(Fig. 48B). Mesepisternum with at least some black hairs (Fig. 48A), often
_	with extensive dark pubescence160
160	Face long, majority of the clypeus passing below an imaginary line drawn
	between the ventral margins of the compound eyes. Clypeus comparatively
	sparsely punctate, punctures separated by an average of 1 puncture diam-
	eter, with impunctate longitudinal midline. Restricted to mountainous ar-
	eas in northern and north-western Spainallosa Warncke (note, the
	Spanish taxon may be distinct from populations in Central Europe)
_	Face short, only a small part of the clypeus falls below an imaginary line
	drawn between the ventral margins of the compound eyes. Clypeus compar-
	atively densely punctate, punctures separated on average by 0.5-1 puncture
	diameter
161	Clypeus extremely densely and coarsely punctate, punctures separated by
	0.5 puncture diameters (Fig. 48B). Currently known only from the Sierra
	Nevada
_	Clypeus less densely punctate, punctures separated on average by 0.5-1
	puncture diameters. Widespread across Iberiabicolor
	Fabricius sensu lato (two mitochondrial lineages are present in <i>A. bicolor</i> ;
	to date, only the southern lineage has been found in Iberia)
162 (151)	Clypeus punctate, interspaces forming weakly raised longitudinal wrinkles
	(Fig. 70E; former subgenus <i>Zonandrena</i>)
_	Clypeus without longitudinal wrinkles
163	Face, mesepisternum, scutum anteriorly, and propodeum with abundant
	white hairs; scutum medially with contrasting short black pubescence.

¹⁵ These two species are very difficult to separate in the female sex. Association with males should be made where possible.

	Marginal areas of 12–4 laterally with broad white hair patches that obscure the underlying surface. Restricted to steppic areas in central Spain
	soror Dours
_	Face with pubescence variable, either black or a mixture of black and brown;
	if entirely white, then mesosoma with extensive brown pubescence dorsally.
	Terga with complete apical hair bands that are never interrupted medially in
	fresh specimens or with hair bands reduced and essentially absent164
164	Facial pubescence white. Rare, restricted to the Pyrenees gravida Imhoff
_	Facial pubescence black or a mixture of black and brown; never pure white
	165
165	Posterior face of hind femur with a latitudinal carina. Facial pubescence
	never entirely black, usually a mixture of yellow, brown, and black hairs,
	sometimes whitish. Terga usually with strong apical hair bands (Fig. 70F;
	beware abraded specimens). The most common and widespread Iberian
	Andrena species
_	Posterior face of hind femur rounded, without a latitudinal carina. Facial
	pubescence dark, often entirely black (Fig. 11C), usually predominant-
	ly black with at most some intermixed dark brown hairs. Terga with hair
	bands variable, sometimes weakly present, sometimes almost entirely absent
	(Figs 11F, 12D). Generally rare and found in southern Iberia166
166	Propodeal triangle with finely raised rugae covering entire surface. Pubes-
	cence variable, from almost entirely melanic (Fig. 70G, with only the tibial
	scopa remaining orange-red) to light, with extensive brown hairs on the
	mesosoma and terga (Fig. 70H). Foveae comparatively weakly constricted
	medially. Usually with complete tergal bands present 16 vulcana Dours
_	Propodeal triangle with finely raised rugae covering only basal half, api-
	cal parts with fine granular shagreenation. Pubescence variable, but nev-
	er entirely melanic, mesonotum always with at least some brown hairs
	(Figs 11D, 12B). Foveae comparatively strongly constricted medially. Tergal
	bands usually greatly reduced 16
167 (162)	Dorsolateral surface of the propodeum reticulate, with large and shal-
	low punctures (can be small in A. nuptialis Pérez), clearly contrasting the
	shagreened and shiny propodeal triangle, this lacking lateral carinae and
	becoming shinier on the declivity (Fig. 65H, often entirely shagreened in
	A. nuptialis). Pronotum lacking lateral keel. Clypeus typically weakly to
	moderately domed, densely punctate (punctures separated by 1 punctures
	diameter, with more or less pronounded impunctate midline (subgenus
	Hoplandrena)
_	Without this combination of characters, pronotum keeled or not173
168	Hind tibiae and/or tarsi golden-orange
_	Both hind tibiae and tarsi dark, at most obscurely reddish-brown170

¹⁶ These two species are difficult to separate due to enormous colour variation.

169	Hind tibiae golden-orange. Clypeus medially with a clearly raised longi-
	tudinal impunctate area, this area smooth and shiny, strongly contrasting
	the remaining parts of the clypeus which are densely punctate. Northern
	and central Spain, in areas with deciduous forest. Associated with Quercus
	(Fagaceae)
_	Hind tibiae dark. Clypeus with faint impunctate longitudinal line medi-
	ally, but this is not raised and not shiny, therefore not strongly contrasting
	with the remaining parts of the clypeus which are regularly punctate. Rare,
	restricted to the Pyrenees
170	A3 clearly exceeding length of A4+5nuptialis Pérez
_	A3 not clearly exceeding length of A4+5, usually as long as or slightly short-
	er than A4+5
171	Metasomal terga with short hairs, most clearly seen in profile on T2-3 with
	hairs not exceeding width of a flagellum rosae Panzer (partim, dark form)
_	Metasomal terga with extensive and abundant long hairs, most clearly seen in
	profile on T2-3 with hairs clearly exceeding width of a flagellum172
172	Tibial scopa in fresh specimens usually dark dorsally and silver ventrally.
	Usually with light brown facial hair. Usually univoltine, flying April to
	mid-June, with potential sporadic emergence in August and September
	(not yet observed in Iberia). Rare, restricted to cooler parts of Iberia 17
_	Tibial scopa in fresh specimens dark dorsally and golden ventrally, but this
	can be ambiguous and fade to silver in older specimens or pinned material.
	Facial hair can be dark, particularly in the spring generation, the summer
	generation usually has lighter facial hair. Bivoltine, usually flying March-
	May and July-August, with phenology depending on local conditions.
	Common and widespread across Iberia 17
	trimmerana (Kirby) (partim, dark form)
173 (167)	Pronotum laterally keeled, angulate, keel runs up dorsally to an angled corner
1/3 (10/)	
_	Pronotum laterally rounded, without a keel205
174	Mesepisternum deeply and distinctly punctate, punctures separated by <1
1/1	puncture diameter, with weakly shiny interspaces. Ocelloccipital distance
	exceeding 3 times the diameter of a lateral ocellus. Terga extremely densely
	punctate, punctures separated by <0.5 puncture diameters, with shiny inter-
	spaces. No recent records, possibly extinct, restricted to central Spain
	Without this combination of characters, mesepisternum usually impunctate
	or only obscurely punctate, ocelloccipital distance shorter, terga dull or less
	densely punctate

¹⁷ These two species cannot be consistently morphologically separated with confidence. Association with males should be made. See discussion and notes in Wood et al. (2022c).

175	Fore margin of clypeus slightly upturned, forming wide] shape, dorsolateral
	surface of propodeum reticulate, impunctate, not strongly differentiated from
	the propodeal triangle, terga with dense apical hair bands on T2-4, fovea broad
	and occupying over ½ of the distance between the compound eye and the
	lateral ocellus, tibial scopae with hairs long and loose. Restricted to central,
	southern, and eastern Spain and associated with <i>Reseda</i> (Resedaceae) 176
_	Without this combination of characters177
176	Terminal fringe orange. Tibial scopae orange. Tergal discs with extremely
	fine shagreen, shiny, clearly and deeply punctate relata Warncke
_	Terminal fringe dark brown medially and white laterally (Fig. 40D). Tibial
	scopae white (Fig. 40A). Tergal discs strongly shagreened, weakly shiny, ob-
	scurely and shallowly punctate (Fig. 40D)blanda Pérez
177	Pygidial plate flat to weakly convex, outer margin usually somewhat ele-
	vated, without clearly raised area medially (Fig. 71A; subgenus Notandrena
	partim)178
_	Pygidial plate with distinctly limited raised area medially (Fig. 71B)186
178	Scutum with very dense and even punctation across the entire surface of
	the disc, punctures separated by much less than the diameter of a puncture,
	in some cases nearly touching (Fig. 57C). Underlying integument strongly
	shagreened and dull griseobalteata Dours
_	Scutum with moderate and uneven punctation, punctures separated by 1–3
	puncture diameters in some cases. If occasional punctures are close to touch-
	ing, then this is not consistently replicated across the whole scutum, with
	other punctures being separated by a clear distance. Underlying integument
	variable, from shagreened to shiny179
179	Hind tibia and all tarsi completely orange. Restricted to areas close to the
	Pyrenees
_	Hind tibia black, tarsi may be black or orange
180	Larger species, body length 11-12 mm. T2-4 with thick white apical hair
	bands that obscure the underlying surface, only weakly interrupted medially
	on T2
_	Smaller species, body length 7–10 mm. T2–4 usually with weaker hair
	bands
181	Tibial scopa, when viewed in reverse profile (i.e. looking at the posterior
	face of the hind tibia) dorsally short and thick, hair length only 1–1.5 times
	the diameter of a lateral ocellus, clearly shorter than the ventral scopal hairs.
	Restricted to areas close to the Pyrenees
_	Tibial scopa equally long dorsally and ventrally
182	Terga very sparsely punctate, punctures of discs of T2–4 shallow, obscure, and
	separated by 3–4 puncture diameters. Clypeus comparatively flattened. Very
	rare, known only from southern Spain (Cádiz)
_	Terga densely punctate, punctures on discs of T2–4 strong and deep, sepa-
	rated at most by 2 puncture diameters, usually by <2 puncture diameters.
	Clypeus comparatively domed

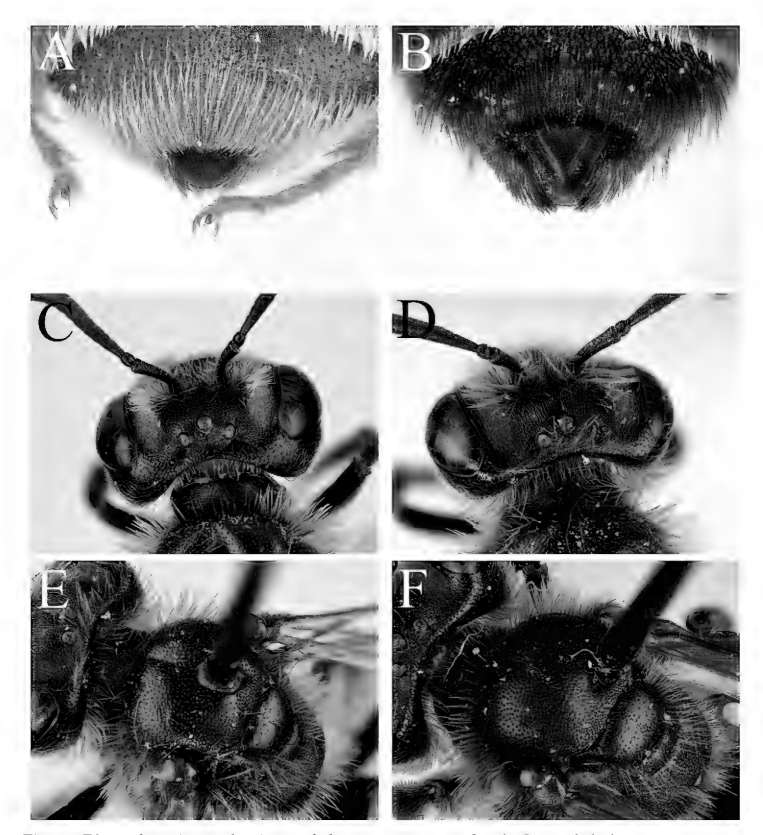


Figure 71. Andrena (Notandrena) griseobalteata Dours, 1872, female **A** pygidial plate, posterior view; Andrena (Leucandrena) leptopyga Pérez, 1895, female **B** pygidial plate, posterior view; Andrena (Notandrena) foeniculae Wood, 2020, female **C** head, dorsal view; Andrena (Notandrena) nitidiuscula Schenck, 1853, female **D** head, dorsal view **E** scutum, dorsolateral view; Andrena (Notandrena) fulvicornis Schenck, 1861, female **F** scutum, dorsolateral view.

184	Central line in the front half of the scutum strongly impressed (Fig. 71E).
	Restricted to temperate habitats in northern Iberia. Hind basitarsi dark.
	Univoltine (July–August)
_	Central line in the front half of the scutum only weakly and superficially im-
	pressed (Fig. 71F). Hind basitarsi dark to often entirely lightened orange. Not
	restricted to northern Iberia, and active in the spring or the summer 185
185	Disc of T1 densely punctate, punctures separated by 1 puncture diameter.
	Scutellum shagreened and dull. Hind basitarsi orange. Throughout Iberia.
	Bivoltine (March–August)
_	Disc of T1 sparsely punctate, punctures separated by 2-3 puncture diam-
	eters. Scutellum polished and shiny. Restricted to strongly saline habitats
	in southern Iberia. Voltinism unclear, possibly bivoltine, recorded March-
	early Junejuliana Wood
186	Flying exclusively in the summer (July–September). Hind tibiae triangular,
	clearly much broader apically than basally, with short scopal hairs not great-
	ly exceeding the diameter of a lateral ocellus (Fig. 72A). Terga with strong
	and broad apical tergal hair bands (Fig. 72B; subgenus <i>Cnemidandrena</i>
	Hedicke)
_	Flying predominantly in the spring, some species extending into July. Hind
	tibiae normal, not greatly broader apically than basally, with long scopal
	hairs, clearly greatly exceeding the diameter of a lateral ocellus. Terga with
	or without clear apical hair bands189
107	•
187	Outer surface of the galea smooth and shiny. Associated with Ericaceae,
	found in Atlantic habitats across northern, central, and western Iberia
	fuscipes (Kirby)
100	Outer surface of the galea shagreened and dull
188	Mesosoma anteriorly and posteriorly with pale hairs, medially with abun-
	dant black hairs. Face with pale hairs. Associated with Asteraceae, restricted
	to northern Spain
_	Mesosoma with pale to brown hairs, with at most occasional intermixed
	black hairs. Face black-haired. Polylectic, though often found on Asteraceae.
	Restricted to areas surrounding the Pyrenees and the Cantabrian Mountains
	with isolated populations in high mountains in southern Spain (particularly
	the Sierra Nevada)
189	Process of the labrum either elongate (as long as or slightly longer than broad)
	or pointed triangular, never medially emarginate (Fig. 72C–H). Clypeus some-
	times with transverse wrinkles (subgenus <i>Leucandrena</i>)190
_	Process of the labrum trapezoidal, always broader than long, usually me-
	dially emarginate, at least weakly. Clypeus never with transverse wrinkles
	(subgenus <i>Andrena</i> s. str.) 197
190	Fovea very wide, occupying the entirety of the space between the inner mar-
	gin of compound eye and the lateral ocellus191
_	Fovea narrower, occupying between half and two-thirds of distance between
	the inner margin of compound eye and the lateral ocellus

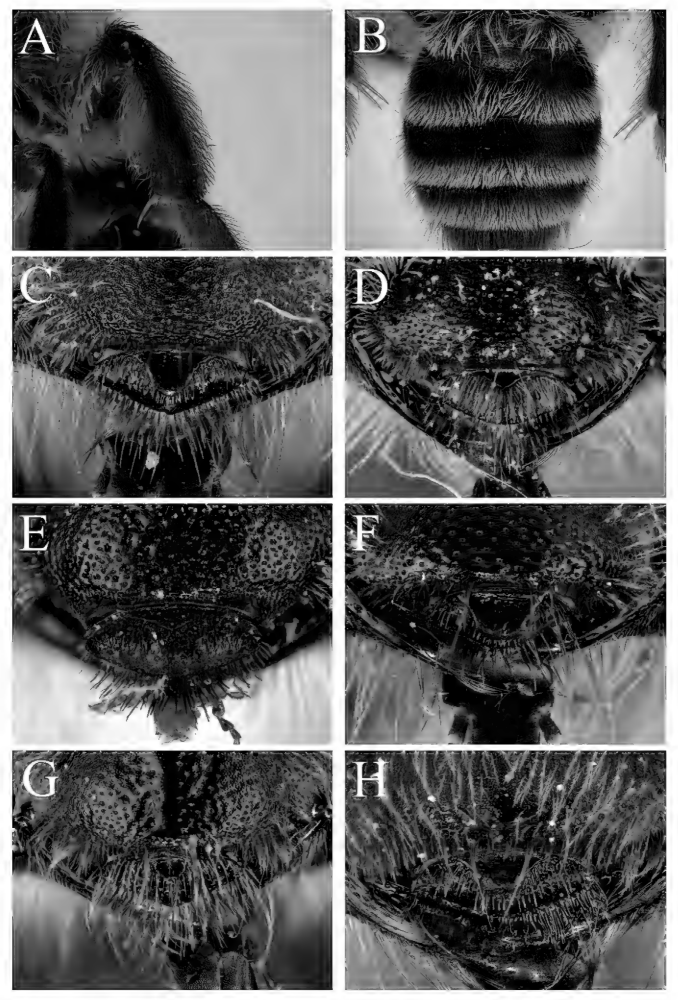


Figure 72. Andrena (Cnemidandrena) denticulata (Kirby, 1802), female **A** hind leg, profile view **B** terga, dorsal view; Andrena (Leucandrena) barbilabris (Kirby, 1802), female **C** process of the labrum, frontal view; Andrena (Leucandrena) tunetana Schmiedeknecht, 1900, female **D** process of the labrum, frontal view; Andrena (Leucandrena) sericata Imhoff, 1868, female **E** process of the labrum, frontal view; Andrena (Leucandrena) parviceps Kriechbaumer, 1873, female **F** process of the labrum, frontal view; Andrena (Leucandrena) ventralis Imhoff, 1832, female **G** process of the labrum, frontal view; Andrena (Leucandrena) leptopyga Pérez, 1895, female **H** process of the labrum, frontal view.

191	Clypeus with network of fine, raised ridges that extend laterally across the disc, underlying surface shiny. Process of labrum short, broader than long, produced
	to a fine triangular point (Fig. 72D)tunetana Schmiedeknecht
_	Clypeus without any lateral ridges, underlying surface shagreened, dull. Pro-
	cess of labrum short and slightly broader than long, but clearly trapezoidal,
	apical margin slightly thickened and raised (Fig. 72H) <i>leptopyga</i> Pérez
192	Scutum and scutellum shagreened, dull. Process of labrum as long as broad,
172	evenly rounded apically (Fig. 72C)
	At least some part of the scutum or scutellum shiny. Process of labrum either
_	,
102	pointed triangular or apically truncate, not evenly rounded apically 193
193	Process of labrum pointed triangular (Fig. 72E). Very rare, restricted to the
	Pyrenees sericata Imhoff
104	Process of labrum apically truncate, not pointed (Fig. 72F, G)
194	Terga clearly finely and densely punctured, punctures separated by 1–2 punc-
	ture diameters, visible against weakly shagreened integument. Restricted to
	mountainous areas of central and northern Spainargentata Smith
_	Terga with punctures sparse and difficult to see, separated by 3–5 puncture
	diameters
195	Process of labrum comparatively large, apex more rounded (Fig. 72F).
	Fovea, viewed dorsally, with dark brown hairs. Terminal fringe dark brown.
	Larger, 10–11 mm
_	Process of labrum comparatively small, more clearly truncate (Fig. 72G).
	Fovea, viewed dorsally, with light brown hairs. Terminal fringe golden.
	Smaller, 7–9 mm
196	Scutum shagreened laterally, but centrally shagreenation is absent, underly-
	ing surface therefore smooth and shiny. Lateral faces of the propodeum with
	clearly raised more or less parallel ridges. Scutum more strongly punctured,
	individual punctures larger and closer together, particularly anteriorly where
	they are separated by 1–2 puncture diameters
_	Scutum shagreened, shagreenation weaker centrally but still clearly visible.
	Lateral faces of the propodeum with at most very small wrinkles. Scutum
	less strongly punctured, individual punctures normal and more scattered,
	separated by 1–3 puncture diameters
197	Terga densely covered with long hairs, in fresh specimens these obscuring
	the underlying surface (Fig. 73A)
_	Terga less thickly haired, sometimes with hair tufts on T1-2, but these not
	obscuring the underlying surface (Fig. 73B)199
198	Hind tibiae orange. Terga predominantly black-haired, sometimes with
	light brown hairs on T1. Restricted to northern Spain, associated with Salix
	(Salicaceae)
_	Hind tibiae dark. Terga 1-5 with extensive reddish-orange pubescence
	(Fig. 73A). With a patchy distribution from northern Portugal and Spain
	to the Pyrenees, with an isolated population in the Sierra de Cazorla (Jáen).
	Associated with various flowering trees and shrubs

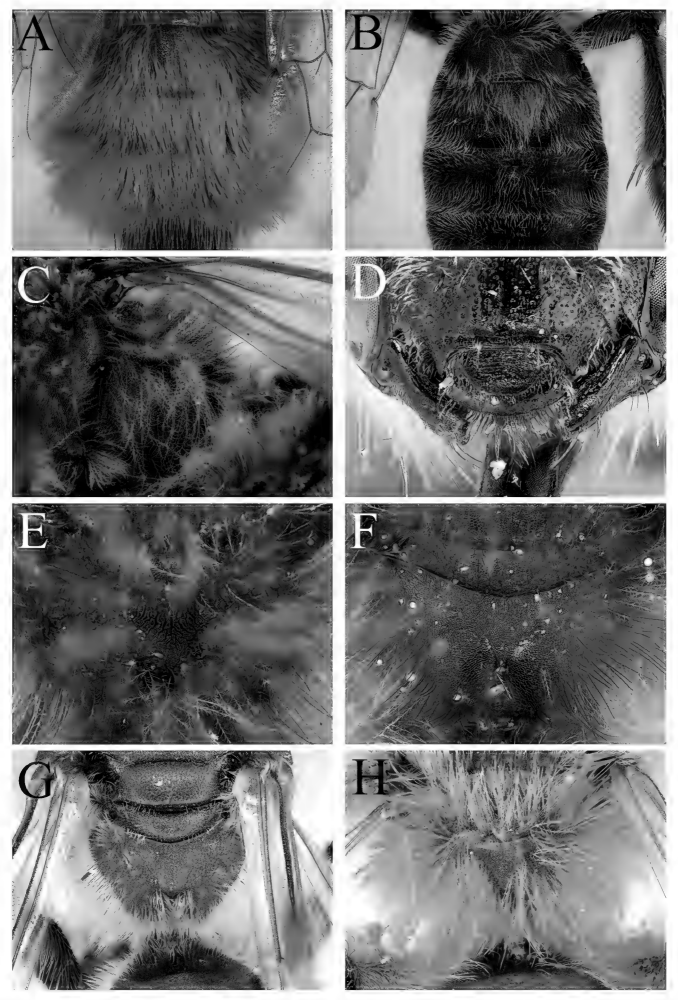


Figure 73. Andrena (Andrena) fulva (Müller, 1776), female A terga, dorsal view; Andrena (Andrena) helvola (Linnaeus, 1758), female B terga, dorsal view; Andrena (?Euandrena) ramosa Wood, 2022, female C propodeal corbicula, profile view; Andrena (Didonia) mucida Kriechbaumer, 1873, female D process of the labrum, ventral view; Andrena (Avandrena) erodiorum Wood & Ortiz-Sánchez, 2022, female E propodeum, dorsal view; Andrena (Avandrena) melacana Warncke, 1967, female F propodeum, dorsal view; Andrena (Nobandrena) funerea Warncke, 1975, female G propodeum, dorsal view; Andrena (Truncandrena) ferrugineicrus Dours, 1872, female H propodeum, dorsal view.

199	Marginal areas of T2–4 very wide, occupying ¾ of each segment
_	Marginal areas of T2–4 never occupying more than ½ of each segment, usually covering only ⅓
200	Face with entirely pale hairs, at most with a few scattered dark hairs along
200	the inner margin of the compound eye
_	Face with extensive black hairs, particularly around the antennal insertions
	and along the inner margin of the compound eye203
201	Terga sparsely haired, with at most weak hair tufts on T1-2. Terga finely
	shagreened and thus weakly shiny. Restricted to northern Spain. Flying
	later in the year (June–August), associated with shrubs, particularly <i>Rubus</i>
	(Rosaceae)
_	Terga typically with long hairs, in fresh specimens with clear and dense
	hair tufts on T1–2. Abraded or older specimens may lack such hair tufts,
	in which case use the following characters: terga strongly shagreened, dull.
	Flying earlier in the year (March–May), associated with flowering trees
202	Resiters of hind tibies parallel sided not conversing anicelly. Destricted to
202	Basitarsi of hind tibiae parallel-sided, not converging apically. Restricted to the Pyrenees with an isolated population in the Sierra de Cazorla (Jáen). As-
	sociated with various flowering trees and shrubshelvola (Linnaeus)
	Basitarsi of hind tibiae broader basally, narrower apically, therefore con-
_	verging apically. Restricted to areas around the Pyrenees and the Canta-
	brian Mountains. Associated with <i>Salix</i> (Salicaceae)
203	Face entirely black haired, without any pale hairs. Terga basally (T1–2) with
_00	orange-brown hairs, apically (T3–4) with extensive and strongly contrast-
	ing black hairs. Associated with <i>Vaccinium</i> (Ericaceae)
_	Face with at least some pale hairs, particularly around the antennal inser-
	tions. Terga without extensive areas with black hairs, generally with mixture
	of predominantly yellowish to brownish hairs on T1-4. Species associated
	with Salix (Salicaceae)
204	Smaller, 10-11 mm. Clypeus predominantly shagreened and dull, with only
	the narrow longitudinal impunctate midline weakly shiny. Terminal fringe
	dark brown
_	Larger, 11-14 mm. Clypeus comparatively shinier, only laterally sha-
	greened and dull, apico-medially broadly shiny around the comparatively
	broader longitudinal impunctate midline. Terminal fringe black
205 (173)	Large species (over 12 mm in length). With abundant black, brown, and/
	or white pubescence. Clypeus strongly domed. Ocelloccipital distance
	long, at least 2 times the diameter of a lateral ocellus (subgenus <i>Melandrena</i>
	partim)
_	Without this combination of characters; remaining species212

206	Tergal discs T1-3(4) with upstanding mixture of short pale whitish to light
	brown hairs extending over both disc and marginal areas. Tibial scopae
	orange-red, or bicoloured and dorsally dark, ventrally orange (dark form,
	active February-April). Terga sometimes with greasy-bronzy metallic reflec-
	tions
_	Tergal discs never with this sort of pubescence, sometimes with white to brown-
	ish pubescence in basolateral corners of tergal discs, never extending onto mar-
	ginal areas. Tibial scopae entirely black or a combination of black and white,
	never orange-red. Terga dark, without metallic reflections
207	Mesosoma with black and white hairs, never with brown hairs208
_	Mesosoma with black, brown, and/or pale hairs, never with only black and
	white hairs209
208	T2-4 laterally with thick apical patches of white hairs, these strongly con-
	trasting the black integument
_	T2–4 laterally without white hair patches, entirely dark <i>morio</i> Brullé
209	Tibial scopae bicoloured, black dorsally and white ventrally. Mesosoma dor-
20)	sally with bright orange-brown pubescence, laterally with pale pubescence,
	never with black hairs on the mesepisternum. Univoltine, flying April-June.
	Restricted to temperate parts of northern Spain
_	Tibial scopae entirely black. Mesosoma dorsally usually with the pubescence
	darker, laterally with pubescence never pale, usually with abundant black
210	hairs, at most with pubescence of the mesepisternum brown
210	Disc of T1 shagreened and barely punctate, punctures scattered and obscure
	against the underlying sculpture
_	Disc of T1 polished and shiny, strongly punctate, punctures clearly visible
	against the underlying sculpture211
211	Disc of T1 with punctures comparatively sparse, separated by 2 puncture
	diameters thoracica (Kirby)
_	Disc of T1 with punctures dense, punctures separated by up to 1 puncture
	diameter, often separated by less
212 (205)	Propodeal corbicula complete, with both anterior and dorsal fringe, fringes
	composed of long, dense, and extremely plumose yellowish-brown hairs, these
	plumose hairs present also on the mesepisternum, the flocculus, and the femo-
	ral scopae (Fig. 73C), but hairs of the tibial scopae simple (see also illustrations
	in Wood et al. 2022). Known only from south-western Spain (Cádiz, Sevilla),
	flying in the very early spring (January-March) ramosa Wood
_	Propodeal corbicula simple, composed of simple or weakly plumose hairs,
	but if plumose then these not extending onto the mesepisternum, flocculus,
	and femoral scopae213
213	Process of the labrum large, as long as broad, apically rounded, ventral sur-
	face covered with latitudinal wrinkles (Fig. 73D). Fovea narrow, occupying
	slightly less than half the space between the compound eye and the lateral

	ocellus. Associated with <i>Muscari</i> (Asparagaceae)
_	Process of the labrum different, either wider than long, trapezoidal, or api-
	cally pointed, ventral surface never covered with latitudinal wrinkles. Foveae
	narrow or broad
214	Head short and broad, at least 1.3 times wider than long. Fovea short and
	broad, only slightly longer than wide. Small bees, 8–10 mm, with long white
	pubescence in fresh specimens. Associated with <i>Erodium</i> (Geraniaceae) 215
_	Without this combination of characters, usually larger. Never associated
	with <i>Erodium</i>
215	
215	Propodeum and mesepisternum entirely microreticulate, without strong or
	weak raised network of reticulation (Fig. 73F). Mesepisternum laterally with
	extensive intermixed black and white hairs. Terga with sparse and short pu-
	bescence, T2-4 with only weak apical hair bands that do not obscure the
	underlying surface. Known from across southern Spain (Albacete, Cádiz,
	Granada, Málaga) melacana Warncke
_	Propodeum and mesepisternum shiny or with granular microreticulation,
	overlain by strong or weak network of raised rugosity (Figs 47E, 73E).
	Mesepisternum predominantly pale-haired, at most with 30% of hairs
	black. Terga with long erect white hairs, forming dense apical hair bands on
	T2-4 which obscure the underlying surface (Fig. 47F). Known from either
	south-eastern (Albaceae) or south-western (Cádiz) Spain216
216	Propodeum (including propodeal triangle) and mesepisternum with
	strongly produced but fine interlinked network of raised rugosity
	(Fig. 73E). Facial foveae occupying ¾ of space between the compound eye
	and a lateral ocellus. Apical fringe of T5 and hairs flanking pygidial plate
	golden-brown. Known from south-eastern Spain (Albacete)
	erodiorum Wood & Ortiz-Sánchez
_	Propodeum with fine granular shagreen, with weak network of raised ru-
	gosity, propodeal triangle slightly depressed, basal 2/3rds with raised lon-
	gitudinal rugae (Fig. 47E). Facial foveae occupying ½ space between the
	compound eye and a lateral ocellus (Fig. 47C). Apical fringe of T5 and hairs
	flanking pygidial plate dark brown (Fig. 47F). Known from south-western
017	Spain (Cádiz)
217	Terga shagreened and dull, impunctate or with obscure and scattered punc-
	tures (Figs 30D, 39D, 51C)218
_	Terga polished and shiny, at most finely shagreened, clearly and densely
	punctate
218	Propodeal triangle with internal surface covered with fine network of raised
	reticulation219
_	Propodeal triangle with internal surface smooth, entirely granularly sha-
	greened, without raised reticulation (Fig. 73G, H)220

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219	Mid and hind basitarsi dark. Tergal discs in profile view with sparse and short dark hairs, apically with white hair bands. Throughout Iberia
_	Mid and hind basitarsi orange (Fig. 39B). Tergal discs in profile view laterally with orange hair, apically with long yellowish hair bands (Fig. 39D). Restricted to high altitude sites (>1200 m) in the Pyrenees
220	Propodeum with comparatively few hairs, these largely restricted laterally to the dorsal fringe of the propodeal corbicula, propodeum dorsally with sparse pubescence (Fig. 73G). Process of the labrum narrow, only slightly broader than long, more or less triangular with a rounded apex. Dorsolateral surfaces of the propodeum obscurely and shallowly punctate. T2–4 with narrow white apical hair bands. Terminal fringe bright orange. Restricted to central Spain.
	Propodeum covered with long abundant pubescence, no meaningful difference between hairs of the dorsal fringe of the propodeal corbicula and those on the dorsal surface of the propodeum (Fig. 73H). Process of the labrum usually trapezoidal, broader than long, with or without an emarginate front margin. If unclear, then dorsolateral surfaces of the propodeum clearly and coarsely punctate. T2–4 with or without hair bands. Terminal fringe darker, from light brown to dark brown or almost black (subgenus <i>Truncandrena</i>)
221	T2–4 with clear apical hair bands222
_	Terga without apical hair bands, hairs may be present on the tergal discs 224
222	Fovea relatively narrow, occupying ½ the space between the compound eye and the lateral ocellus. Process of the labrum triangular, slightly truncate. Smaller, not exceeding 10 mm in length. Associated with Brassicaceae
_	Fovea broad, occupying ³ / ₄ of space between the compound eye and the lateral ocellus. Process of the labrum trapezoidal with a strong emargination medially. Larger, exceeding 12 mm in length. Associated with <i>Cistus</i> (Cistaceae)
223	Scopa bicoloured, black dorsally and orange ventrally (Fig. 51D). Face longer, clypeus ventrally projecting well below a line drawn between the lower margins of the compound eyes (Fig. 51B). Larger, 15–16 mm. Known only

Mid and hind basitarsi dark, at most obscurely dark reddish.....226

225	Mesepisternum laterally with abundant dark hairs. Base of discs of T2–4
	strongly depressed (depressed relative to the apical margin of the preceding
	tergum, often with a physical space between them), this depression therefore
	laterally exaggerating the gradulus along its inner margin. Tergal margins
	(and sometimes sides of terga when viewed laterally) sometimes lightened
	reddish, contrasting the dark discs. Rare, known only from southern Spain
	(Alicante, Cádiz)
_	Mesepisternum laterally with entirely pale hairs. Base of T2-4 not noticea-
	bly depressed, therefore gradulus not particularly noticeable. Tergal margins
	always dark. Common throughout Iberia
226	Discs of T2–4 with abundant and extensive pale pubescence, in fresh speci-
	mens this pubescence forming distinct patches laterally. Marginal areas broad
	on T3–4 occupying $^2/_5$ of the length of the segment. Clypeus shagreened and
	dull, with a narrow impunctate longitudinal mid line, only slightly shiny at
	extreme apex. Throughout Iberia
_	Discs of T2–4 with at most scattered and fine hairs, never forming patches
	Marginal areas narrow, on T3–4 occupying at most 1/5 th of the length of the
	segment. Clypeus less extensively shagreened and dull, becoming smooth and
	shiny in its apical half, with broader impunctate longitudinal mid line. Rare
	known only from southern Spain (Córdoba)
227	Terga without apical hair bands, or with obscure hair bands that are widely
22/	interrupted medially even in fresh specimens (Fig. 74A, B). Tibial scopae
	bicoloured, dark dorsally and white ventrally. Terminal fringe dark brown
	to black, with some white hairs laterally228
	Terga with clear apical hair bands, these complete at least on T3—4
_	(Fig. 74C, E, F). Tibial scopae unicolourous white or golden-orange. Ter-
	minal fringe light, reddish-brown to golden229
228	Terga with widely interrupted apical hair bands, often abraded and absent
220	
	Terga coarsely and densely punctate, tergal margins strongly depressed, with
	punctures at most half the size of those on the tergal discs (Fig. 74A). Rare
	restricted to southern Iberia
_	Terga without apical hair bands. Terga extremely finely punctate, no major
	difference in the size of the punctures on the weakly depressed marginal
	areas compared to the discs (Fig. 74B). More widespread across southern
	central, and eastern Iberiacorax Warncke
229	Scutum sparsely punctate, punctures separated by 1–2 puncture diameters
	laterally shagreened and dull, shagreenation weakening medially and or
	the scutellum, here shiny. Process of the labrum short and triangular (Fig.
	74D)urdula Warncke
_	Scutum and scutellum densely punctate, punctures separated by up to 1 punc-
	ture diameter, underlying surface uniformly finely shagreened and weakly shiny
	Process of the labrum trapezoidal, margin truncate to weakly emarginate 230

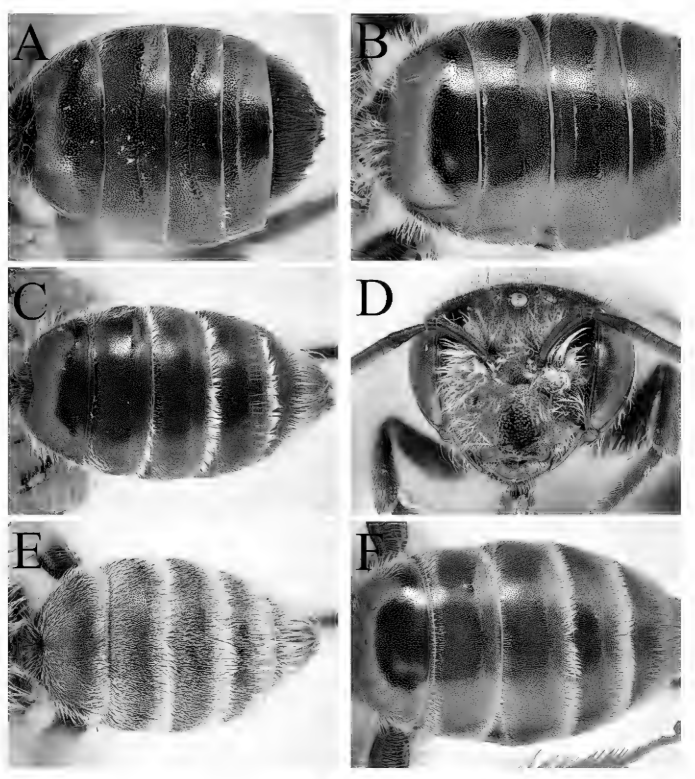


Figure 74. Andrena (incertae sedis) macroptera Warncke, 1974, female **A** terga, dorsal view; Andrena (incertae sedis) corax Warncke, 1975, female **B** terga, dorsal view; Andrena (incertae sedis) urdula Warncke, 1965, female **C** terga, dorsal view **D** face, frontal view; Andrena (Ovandrena) farinosa Pérez, 1895, female **E** terga, dorsal view; Andrena (incertae sedis) murana Warncke, 1975, female **F** terga, dorsal view.

For the male key, the following shortcuts can be used:

A.	Clypeus at least partly yellow-marked
В.	Propodeal triangle clearly defined by strongly raised carinae, internal surface rugose-areolate
C.	Genital capsule distinctive, with clear 90° emargination in the outer parts of the gonostyli. The most common <i>Andrena</i> species in Iberia
	go to Andrena flavipes Panzer
D.	Process of the labrum thickened and expanded, anteriorly projecting beyond the fore margin of the clypeus. Pronotum with strong humeral angle. Active in the summer (mid-June to September) (subgenus <i>Cnemidandrena</i> and remaining <i>Margandrena</i>)
Е.	Fore margin of clypeus upturned AND gena conspicuously broadened, wider than the width of the compound eye AND pronotum with a conspicuous humeral angle (remaining <i>Notandrena</i>)
F.	Small species, body length of 5–7 mm (exceptionally 8 mm), terga always dark (remaining <i>Aciandrena</i> , <i>Aenandrena partim</i> , <i>Avandrena</i> , <i>Cordandrena</i> , <i>Cryptandrena partim</i> , <i>Graecandrena</i> , and almost all <i>Micrandrena</i>)
G.	Mesepisternum and/or dorsolateral parts of the propodeum conspicuously punctate AND ocelloccipital distance at least 3 times the diameter of a lateral ocellus (<i>incisa</i> -group, <i>Pruinosandrena</i>)
H.	Clypeus flattened over majority of its surface (subgenus <i>Taeniandrena</i>)
I. J. K.	At least some tergal discs red-marked
L. M.	eye
N.	Remaining species; measured along ventral margin, A3 slightly longer than A4
Male	es es
1	Clypeus at least partly yellow-marked (Figs 44D, 75A–F); marking may be reduced to small marking medially or apically2
_ 2	Clypeus entirely dark, never with yellow markings
_	Yellow facial markings restricted to the clypeus, paraocular areas entirely dark (Figs 44D, 75D–F)
3	Propodeum almost entirely declivous, without clearly differentiated horizontal and vertical parts. Propodeum with dorsolateral parts adjacent to

	the propodeal triangle densely and deeply punctate, punctures separated by
	<0.5 puncture diameters (c.f. Fig. 65C)
_	Propodeum with clearly differentiated horizontal and vertical parts. Propodeum with an exist part of his parts with a resistant by
	deum with or without punctures, but never with punctures separated by
,	<0.5 puncture diameters
4	Ocelloccipital distance 2.5–3 times the diameter of a lateral ocellus. Tergal
	discs occasionally red-marked
_	Ocelloccipital distance <2 times the diameter of a lateral ocellus. Tergal discs always dark
5	S5 apically with extremely dense and pronounced latitudinal tuft of yellowish hairs. Terga with apical hair bands broadly interrupted. Typically flying earlier, May-June
_	S5 apically without dense latitudinal hair tuft, at most with scattered hairs. Terga typically with clear and uninterrupted apical hair bands, though beware abraded specimens. Typically flying later, July-August decipiens Schenck
6	Discs of T2–3 entirely and conspicuously red-marked
_	Discs of T2–3 dark, at most with the apical margins lightened8
7	Large species, 11–13 mm. Mandibles long, sickle-like, crossing in their apical third (Fig. 74A). Head short, clearly broader than long. Genital capsule with gonostyli broad, weakly converging apically (Fig. 75G)schencki Morawitz
_	Smaller species, never exceeding 9 mm. Mandibles normal, not strongly crossing apically. Head more or less round. Genital capsule with gonostyli apically narrow, strongly converging subapically (Fig. 75H)
8	Paraocular areas with markings narrow, running up the inner margin of the compound eye, clearly dorsally exceeding the antennal insertions (Fig. 75C). S8 with a ventrally projecting spine at each lateral margin
	orbitalis Morawitz
_	Paraocular areas with markings more or less rectangular or quadrangular, never narrow and running up the inner margin of the compound eye (Fig. 75B); typically not exceeding the antennal insertions dorsally9
9	Fore margin of the clypeus clearly and strongly upturned. Pronotum with strong carinate humeral angle. Head broad, clearly broader than long10
_	Fore margin of the clypeus normal, not upturned. Pronotum without a strong humeral angle. Head variable, but not strongly broadened13
10	Scutum strongly and densely punctate, punctures separated by 0.5–1 puncture diameter, underlying surface shagreened. A5 long, clearly exceeding the length of A4; A4 at most 0.6 times as long as A5. Larger, 9–10 mm
	Scutum either strongly sharreened and obscurely punctate or with shiny
_	Scutum either strongly shagreened and obscurely punctate, or with shiny interspaces, never strongly punctured and shagreened. A5 short, not noticeably longer than A4, both segments quadrate or subquadrate. Smaller, 8–9 mm

11	Gena posteriorly rounded, without carina. All tarsi and the apex of the hind tibia lightened orange. Scutum strongly shagreened and obscurely punctate.
	Flying April-June
_	punctures12
12	Hind tarsi dark. Genital capsule with penis valves comparatively narrow.
12	Associated with saline soils in southern Iberia. Flying March-June
	juliana Wood (partim, with yellow markings on lower paraocular areas)
_	Hind tarsi lightened orange. Genital capsule with penis valves broad, oc-
	cupying the majority of the space between the gonostyli. Restricted to the
	Pyrenees. Flying July-August
13	Terga clearly and finely shagreened, dull to weakly shiny, with small and weak
	punctures. Facial markings yellow. Ocelloccipital distance short, equalling the
	diameter of a lateral ocellus. Restricted to central Spain funerea Warncke
_	Terga smooth and shiny between strongly pronounced punctures. Facial
	markings white. Ocelloccipital distance larger, at least 2 times the diameter
	of a lateral ocellus14
14	Larger, exceeding 10 mm in length. Mandibles basally with white markings.
	Body with abundant white pubescence, with dense hairs on the clypeus and
	ventrally on the gena that can obscure the underlying surface in fresh individu-
	als. Widespread across all of Iberia. Flying April–Julyleucolippa Pérez
_	Small species, 7–8 mm. Mandibles dark, without basal white markings
	(Fig. 75B). Body with reduced pubescence, never with dense hairs that ob-
	scure the underlying surface. Rare, restricted to cool temperate habitats in
	and around the Pyrenees and Cantabrian Mountains. Flying July–August
15	Discs of T2–3 clearly and extensively red-marked
1)	Tergal discs dark, at most with the apical margins lightened
16	Clypeus with yellow marking small, occupying only a small proportion of
10	the clypeus medially, without internal black markings (Fig. 75F). Flying
	early in the year, March-May; widespread throughout Iberia. Associated
	with Asphodelus (Asphodelaceae)sardoa Lepeletier
_	Clypeus almost entirely pale, with two small black markings. Flying later
	in the year, from late May to August; restricted to temperate habitats across
	Spain, including mountains in eastern and south-eastern Spain Associated
	with scabious (former Dipsacaceae = Caprifoliaceae)17
17	Pronotum with strong humeral angle. Fore margin of the clypeus laterally pro-
	duced into two small but distinctly projecting points (Fig. 76A). Smaller, 7-9
	mm. Restricted to areas in and around the Pyreneesmarginata Fabricius
_	Pronotum rounded. Fore margin of the clypeus normal, without lateral pro-
	jections. Larger, 14-16 mm. Restricted to montane grasslands in northern
	and central Spain with isolated populations in the Sierra de Cazorla and
	Sierra Nevada

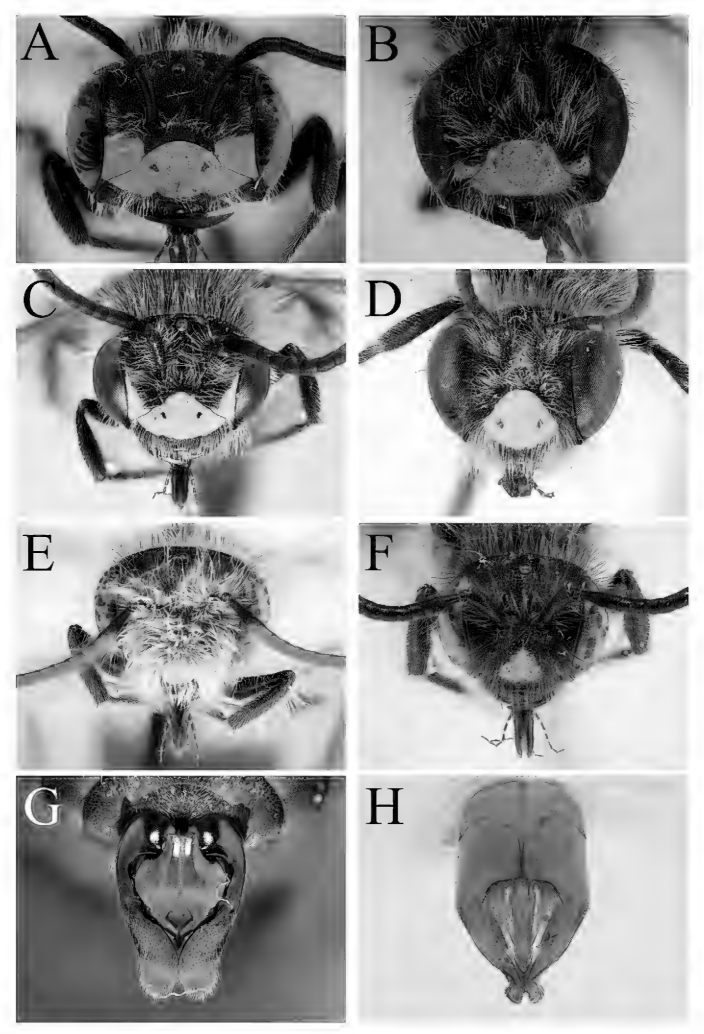


Figure 75. Andrena (Opandrena) schencki Morawitz, 1866, male A face, frontal view G genital capsule, dorsal view; Andrena (Oreomelissa) coitana (Kirby, 1802), male B face, frontal view; Andrena (Rufandrena) orbitalis Morawitz, 1871, male C face, frontal view; Andrena (Orandrena) monilia Warncke, 1975, male D face, frontal view; Andrena (Truncandrena) doursana Dufour, 1853, male E face, frontal view; Andrena (Lepidandrena) sardoa Lepeletier, 1841, male F face, frontal view; Andrena (Poecilandrena) labiata Fabricius, 1781, male H genital capsule, dorsal view.

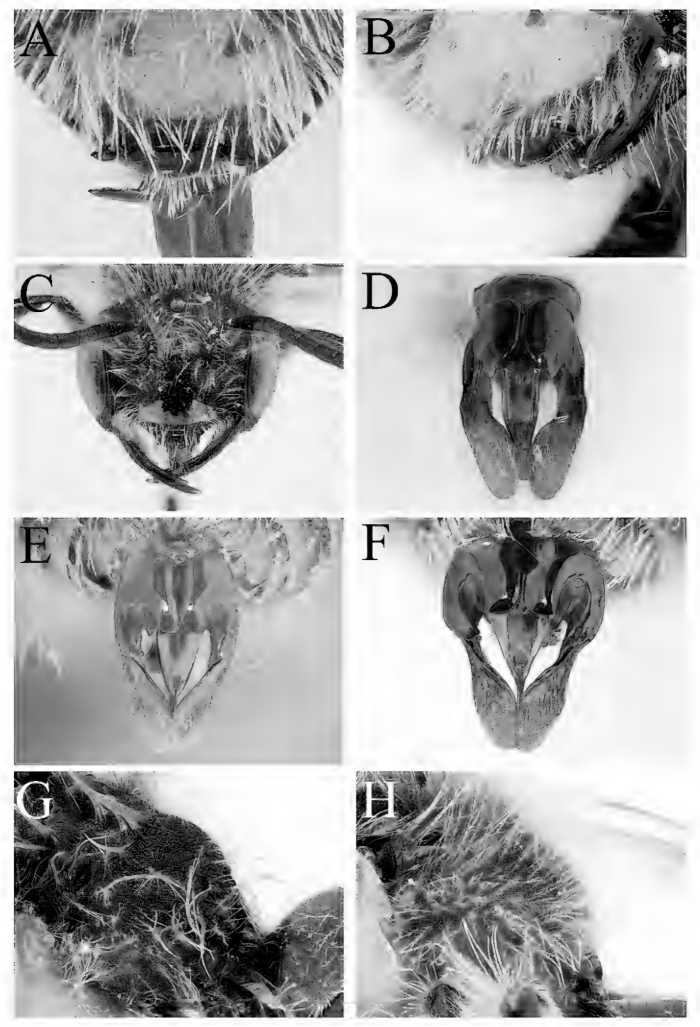


Figure 76. Andrena (Margandrena) marginata Fabricius, 1776, male **A** apex of clypeus, frontal view; Andrena (Holandrena) flavilabris Schenck, 1874, male **B** apex of clypeus, frontolateral view; Andrena (Leucandrena) dinizi Warncke, 1975, male **C** face, frontal view **F** genital capsule, dorsal view **G** propodeum, profile view; Andrena (Leucandrena) tunetana Schmiedeknecht, 1900, male **D** genital capsule, dorsal view; Andrena (Leucandrena) sericata Imhoff, 1868, male **E** genital capsule, dorsal view; Andrena (Leucandrena) ventralis Imhoff, 1832, male **H** propodeum, profile view.

18	Head, mesosoma, and/or metasoma with metallic green or blue reflections
_ 19	Yellow marking of clypeus very small, reduced to a narrow latitudinal strip
	at the fore margin of the clypeus (Fig. 75E)
_	Yellow or white marking of the clypeus always larger, covering the majority or the entirety of the clypeus (with exception of pair of small black dots)20
20	Discs of T2–4 clearly and densely punctate with strong punctures, punctures separated by ≤1 puncture diameter
_	Tergal discs with weaker and more scattered punctures, punctures either fine or separated by >1 puncture diameter
21	Pronotum rounded. Tergal margins not strongly depressed, more or less at the same level as the tergal discs. Gena normal, as wide as the width of the compound eye. Restricted to high elevation sites in the Pyrenees
_	Pronotum laterally carinate. Tergal margins strongly depressed, clearly lower than the level of the tergal discs. Gena broadened, clearly broader than the width of the compound eye. Found in central and southern Iberia
22	Larger, 8–10 mm. Scutum laterally shagreened, medially with a large more or less circular smooth and shiny area, this almost impunctate
	aerinifrons Dours
_	Smaller, 6–7 mm. Scutum uniformly shagreened and punctate, without a smooth and shiny area
23	Inner hind tibial spurs strongly bent at their apexes. Body with subdued metallic reflections, almost without reflections
_	Inner hind tibial spurs straight, parallel-sided. Body with clear metallic re-
	flections
24	Fore margin of the clypeus upturned (Fig. 76B). Pronotum with a strong humeral angle25
_	Fore margin of the clypeus normal, not upturned. Pronotum with or without a humeral angle
25	Propodeum almost entirely declivous, without clearly differentiated horizontal and vertical parts. Propodeum with dorsolateral parts adjacent to the propodeal triangle densely and deeply punctate, punctures separated by <0.5 puncture diameters
_	Propodeum with clearly differentiated horizontal and vertical parts. Propodeum at most obscurely punctate
26	Larger, 10 mm. Scutum and scutellum shagreened, without shiny areas
	langadensis albipila Warncke
_	Smaller, 6–7 mm. Scutum and scutellum at least medially with smooth and shiny areas

27	Gena posteriorly carinate. Surface of gena predominantly smooth and shiny.
	Associated with saline soils in southern Iberia
	juliana Wood (partim, with yellow markings restricted to the clypeus)
_	Gena posteriorly rounded, without carina. Surface of gena microreticulate
	and dull. Very rate, recorded only from the Cádiz region, habitat preference
	unknown microthorax Pérez
28	Mandibles elongate, sickle-like, strongly crossing in their apical third
	(Fig. 76C) AND pronotum with strong humeral angle AND gena
	broadened, clearly broader than the width of the compound eye, usually
	produced into a 90° angulation posteroventrally (yellow-faced <i>Leucan-</i>
	drena)
_	Without this combination of characters; mandibles either shorter, prono-
_	tum laterally rounded, or gena not exceeding width of head
29	,
29	Ocelloccipital distance short, <0.5 times the diameter of a lateral ocellus.
	Genital capsule with gonocoxal teeth apically strongly diverging, apexes
	pointed (Fig. 76D). Found in hot or Mediterranean areas in central and
	southern Iberia with Brassicaceae tunetana Schmiedeknecht
_	Ocelloccipital distance long, 1–2 times the diameter of a lateral ocellus.
	Genital capsule with gonocoxal teeth not strongly diverging, apexes truncate
	(Fig. 76E, F). Found in temperate areas in central and northern Iberia with
	Salix (Salicaceae)30
30	Typically larger, 9–10 mm. Genital capsule more elongate, gonocoxal
	teeth with apexes comparatively narrow, gonostyli not noticeably con-
	stricted medially (Fig. 76E). Very rare, restricted to the Pyrenees
	sericata Imhoff
_	Typically smaller, 7–9 mm. Genital capsule more compact, gonocoxal teeth
	with apexes comparatively broad, gonostyli strongly constricted medially
	(Fig. 76F). More widespread across central and northern Iberia31
31	Lateral faces of the propodeum with a fine network of raised rugosity
0 -	on top of the underlying granular microreticulation (Fig. 76G). Scutum
	with strong and obvious punctures, medially becoming smooth and
	shiny
	Lateral faces of the propodeum with regular granular microreticulation
_	
	(Fig. 76H). Scutum obscurely punctate, uniformly microreticulate and
22	dull
32	Small species, length not exceeding 8 mm
_	Larger species, at least 9 mm in length
33	Propodeal triangle with smooth granular microreticulation, without raised
	rugosity. Very small species, usually not exceeding 6 mm (yellow-faced
	Aciandrena)34
_	Propodeal triangle with at least some raised rugosity, never with uniformly
	smooth microreticulation. Usually a little larger, 6–8 mm35

34	Terga regularly and clearly punctate, punctures extending onto depressed
	tergal margins. Throughout Iberia
_	Terga obscurely punctate, punctures disappearing into underlying microre-
	ticulation, tergal margins more or less impunctate. Restricted to central and
25	eastern Spain
35	Hind basitarsi entirely lightened orange. Frons with contrasting black and
	white pubescence, black on the frons and gena posteriorly, intermixing with
	white around the antennal insertions, becoming white on the clypeus and
	on the gena ventrally. Restricted to <i>Potentilla</i> -rich (Rosaceae) cool habitats
	in the Pyrenees. Flying July-August tarsata Nylander
_	Hind basitarsi dark. Face with uniformly bright white or yellowish pu-
	bescence. Not restricted to the Pyrenees, usually flying earlier in the
26	year
36	Tergal discs covered with upstanding whitish pubescence, this forming dense
	apical hair bands on tergal margins in fresh individuals. Scutum anteriorly sha-
	greened, becoming smooth and shiny posteriorly. Antennae ventrally strongly
	and extensively lightened orange. Restricted to south-western Spain (Huelva,
	Sevilla), flying March-April
_	Tergal discs never with upstanding pubescence; white apical hair bands can be present on tergal margins. Scutum with uniform sculpture, consistently
	shiny across its surface. Antennae typically dark ventrally37
37	Disc of T1 with scattered punctures, punctures separated by 1–3 puncture
37	diameters, strongly contrasting punctation of the discs of T2–4, here punc-
	tures separated by 0.5 puncture diameters. Clypeus yellow-marked across its
	entire surface with the exception of two small black marks. Genital capsule
	compact, with pronounced gonocoxal teeth and small and spatulate gono-
	styli (Fig. 77A). Restricted to central Spain <i>taxana</i> Warncke
_	All tergal discs uniformly punctate, punctures separated by 0.5 puncture
	diameters. Clypeal marking often reduced in size, not covering entire sur-
	face. Genital capsule elongate, with weakly pronounced gonocoxal teeth,
	gonostyli apically produced into triangular wedges (Fig. 77B)
	ventricosa Dours (partim, light form)
38	Mid and hind basitarsi entirely lightened orange (Fig. 44B)39
_	Mid and hind basitarsi dark41
39	Ocelloccipital distance 3 times the diameter of a lateral ocellus. Terga very
	finely shagreened, more or less smooth and shiny. Tergal discs regularly and
	deeply punctate, punctures separated by 1–2 puncture diameters, punctures
	becoming smaller and continuing onto tergal margins. Genital capsule, see
	Fig. 44Hlimbata dusmeti Warncke
_	Ocelloccipital distance <2 times the diameter of a lateral ocellus. Terga
	strongly shagreened, at most weakly shiny, tergal discs with small and obscure
	punctures that disappear into the underlying structure, tergal margins more
	or less impunctate40

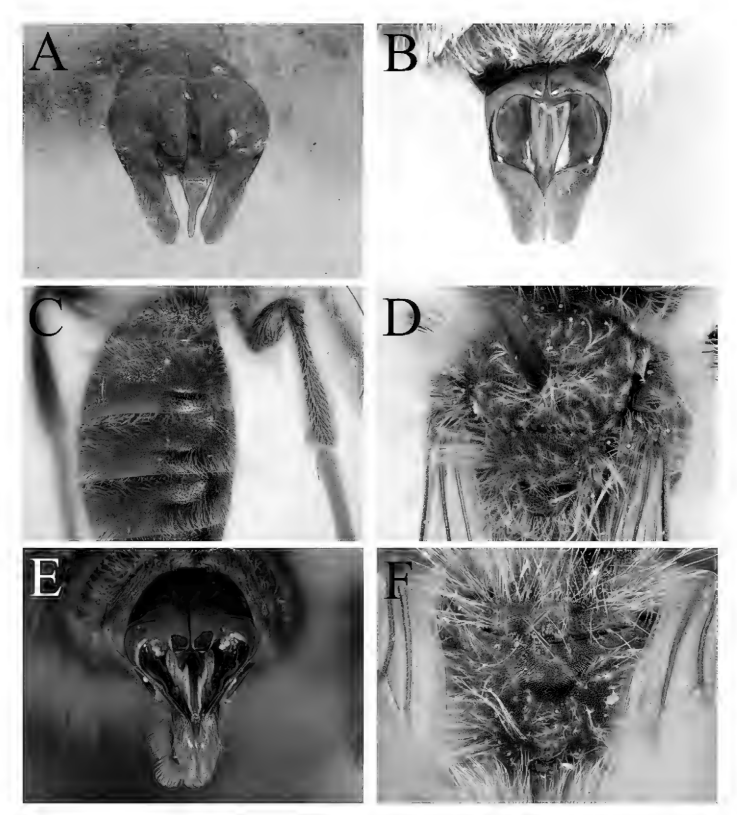


Figure 77. Andrena (Parandrenella) taxana Warncke, 1975, male A genital capsule, dorsal view; Andrena (Cryptandrena) ventricosa Dours, 1873, male B genital capsule, dorsal view; Andrena (Truncandrena) minapalumboi Gribodo, 1894, male C terga, dorsolateral view; Andrena (Chlorandrena) rhyssonota Pérez, 1895, male D scutum and scutellum, dorsal view; Andrena (Charitandrena) hattorfiana (Fabricius, 1775), male E genital capsule, dorsal view; Andrena (Orandrena) monilia Warncke, 1975, male F propodeum, dorsal view.

41	Scutum and scutellum with dense network of raised longitudinal striations, most obvious on the scutellum (Fig. 77D). Tergal margins and tergal bases
	strongly depressed, giving individual tergal segments a domed appearance
	when viewed in profile
_	Scutum and scutellum without longitudinal striations. Tergal margins depressed or not, but not in combination with tergal bases depressed42
42	Large species, 14–16 mm. Genital capsule distinctive, elongate with project-
72	ing and apically truncate gonocoxal teeth (Fig. 77E). Pubescence variable,
	from light brown to almost entirely black (ssp. <i>nigricauda</i> Wood). Restricted
	to montane grasslands in northern and central Spain with isolated popula-
	tions in the Sierra de Cazorla and Sierra Nevada
_	Smaller, sometimes reaching 14 mm, but usually 9–12 mm. Genital capsule
	different; if elongate, then gonocoxal teeth pointed, not truncate. Pubes-
	cence predominantly bright, never almost entirely black
43	Face with strongly contrasting black and white pubescence; clypeus medi-
	ally with white hairs, inner margins of the compound eyes and area around
	antennal insertions and frons with abundant black hairs44
_	Face with bright pubescence, at most with occasional scattered dark hairs,
	never strongly contrasting light pubescence46
44	A3 short, only slightly longer than A4. A4 rectangular, only slightly longer
	than broad, clearly shorter than A5. Antennal segments slightly bulging ven-
	trally. Rare, known only from southern Spain (Córdoba) varia Pérez
_	A3 long, clearly longer than A4. A4 rectangular, clearly longer than broad
	slightly shorter than A5. Antennal segments parallel-sided, without ventral
	bulges
45	Genital capsule with apical flattened part elongate, longer than broad,
	appearing triangular (Fig. 51D, E). Inner margin of these flattened areas
	strongly and acutely raised, slightly reflexed. Larger, 13–14 mm. Known
	only from southern Spain (Málaga)ghisbaini sp. nov.
_	Genital capsule with apical flattened part rounded, as long as broad
	(Fig. 51F). Inner margin of these flattened areas less strongly raised and not
	reflexed. Smaller, 10–11 mm. Throughout Iberiavillipes Pérez
46	Propodeal triangle broad, internal surface with very fine granular reticu-
	lation, laterally and posteriorly weakly shiny, contrasting the dorsolateral
	parts of the propodeum (Fig. 77F). Face in frontal view with compound
	eyes large, almost bulbous, separated by a distance only not greatly
	exceeding the diameter of an individual compound eye (Fig. 75D).
	Genital capsule elongate (Fig. 78A). Restricted to hot steppe in central
	Spain
_	Propodeal triangle with clearer structure of raised reticulation or rugosity,
	without weakly shiny margins. Face in frontal view with compound eyes
	normal, separated by a distance much greater than the diameter of an indi-
	vidual compound eye (Fig. 78B). Genital capsule otherwise
	rigual compound eve (1 iz. / OD). Ciciliai capquic offici wist

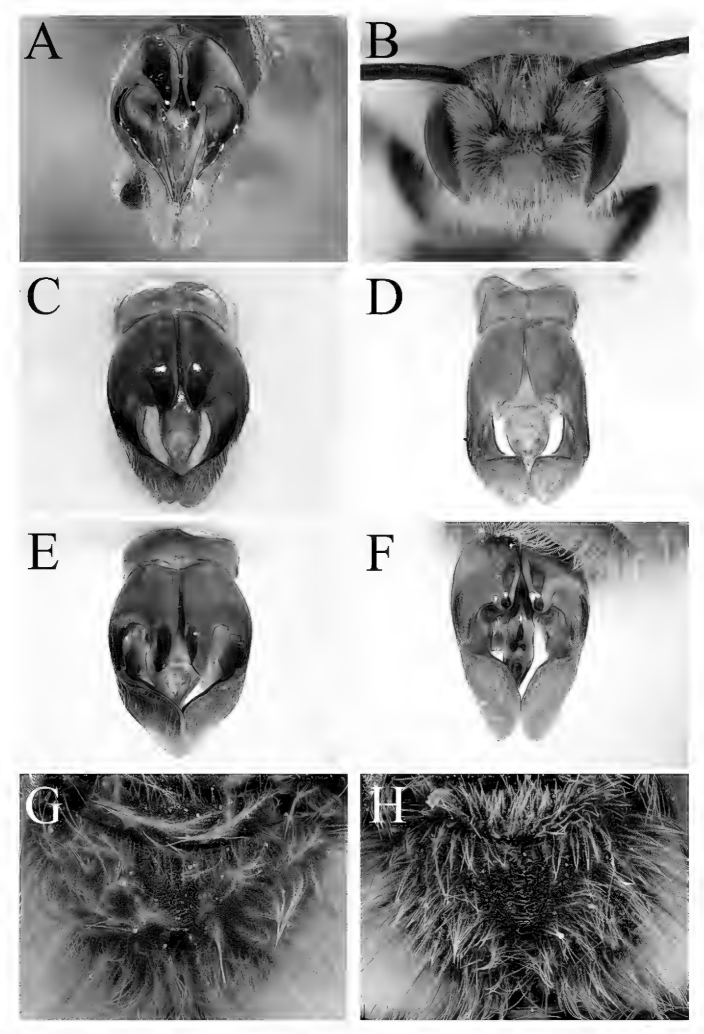


Figure 78. Andrena (Orandrena) monilia Warncke, 1975, male A genital capsule, dorsal view; Andrena (incertae sedis) murana Warncke, 1975, male B face, frontal view D genital capsule, dorsal view; Andrena (incertae sedis) relata Warncke, 1975, male C genital capsule, dorsal view; Andrena (Chlorandrena) humilis Imhoff, 1832, male E genital capsule, dorsal view; Andrena (Truncandrena) nigropilosa Warncke, 1967, male F genital capsule, dorsal view; Andrena (Suandrena) suerinensis Friese, 1884, male G propodeum, dorsal view; Andrena (Plastandrena) pilipes Fabricius, 1781, male H propodeum, dorsal view.

47	Terga finely shagreened, more or less smooth and shiny, regularly and deeply punctate, punctures separated by 1–2 puncture diameters, punctures extending onto the tergal margins. A3 comparatively short, only slightly ex-
_	ceeding length of A4. Nervulus antefurcal
48	cal
_	Genital capsule complex, with inner margins progressively diverging, forming 90° inner angle. Gonostyli apically strongly flattened and broadened, more or less triangular shovel-like (Fig. 78D)
49	Disc of T1 with large 'crater punctures', punctures with distinctly raised margins. Terga shagreened and weakly shiny. Genital capsule with long, strongly produced gonocoxal teeth (Fig. 78E)
_	Disc of T1 with small hair-bearing punctures, without raised, crater-like rims. Terga microreticulate and dull. Genital capsule with only weakly produced gonocoxal teeth
50	A3 very long, slightly exceeding the length of A4+5. A4 very short, broader than long. Genital capsule relatively compact, with flattened apical part
_	A3 comparatively shorter, at most equalling the length of A4+5. A4 longer, slightly longer than broad. Genital capsule relatively elongate, with flattened apical part only slightly longer than broad (Fig. 78F)
51 (1)	Propodeal triangle clearly defined by strongly raised carinae, internal surface rugose-areolate (Figs 78G, H, 79A)
_	Propodeal triangle not strongly defined by lateral carinae with its internal surface rugose-areolate
52	Forewing with two submarginal cells. Clypeus with longitudinal striations. lagopus Latreille
_	Forewing with three submarginal cells. Clypeus without longitudinal striations
53	Small species, 6–7 mm. Clypeus, scutum, and terga extremely densely and deeply punctate with small punctures, punctures almost confluent. Each side of T2 laterally with a small but deep and clearly defined fovea (Fig. 79B, C; subgenus <i>Brachyandrena</i>)
_	Larger species, at least 8 mm in length, usually >10 mm. Body never so densely punctate, punctures always a little separated, never confluent. T2 with fovea obscure

54	T2 laterally with foveae narrow and relatively elongate (Fig. 79B). Common
	and widespread across Iberia 18
_	T2 laterally with foveae shorter and relatively broad (Fig. 79C). Rare, encountered predominantly in southern Spain (Cádiz, Málaga) ¹⁸ <i>miegiella</i> Dours
55	Pronotum laterally with strong humeral angle with well-developed carina.
	Fore margin of clypeus weakly upturned
_	Pronotum laterally rounded. Fore margin of clypeus normal, not upturned
56	Hind basitarsi and majority of hind tibiae lightened orange. Body with in-
<i>)</i> 0	tegument uniformly dark. Pubescence bright, face and mesepisternum with
	light brown hairs, scutum and T6 with bright orange hairs. A3 clearly short-
	er than A4
_	Without this combination of characters; either hind legs dark, body with
	darker pubescence, or A3 clearly longer than A4
57	Terga with metallic reflections and at most superficial punctures AND geni-
71	tal capsule distinctive, with strongly produced gonocoxal teeth and gonosty-
	li strongly reflexed, with long plumose hairs present on their outer margin
	(subgenus Suandrena)58
_	Terga without metallic reflections OR if with metallic reflections, then geni-
	tal capsule otherwise, either lacking pronounced gonocoxal teeth or without
	plumose hairs on the outer margin of the gonostyli OR terga clearly and
	abundantly punctate (subgenus <i>Plastandrena</i>)60
58	Genital capsule with penis valves grossly inflated, occupying entirety of
	space between the gonostyli, laterally produced into bulbous projections
	(Fig. 79D). Known only from Cádiz province, flying in December
	gades Wood & Ortiz-Sánchez
_	Genital capsule with penis valves narrower, essentially parallel-sided, not
	occupying entirety of space between the gonostyli, without lateral projec-
	tions
59	A3 ventrally rounded, lacking any kind of triangular point (Fig. 79E). Geni-
	tal capsule less elongate, gonocoxal teeth relatively short (Fig. 79F). Con-
	firmed males known only from eastern and southern Spain. Bivoltine, flying
	February-April and September-October
_	A3 ventrally produced into triangular point (Fig. 79G). Genital cap-
	sule more elongate, with long and strongly produced gonocoxal teeth
	(Fig. 79H). Widespread across Iberia. Univoltine, flying March-May
_	suerinensis Friese
60	Genital capsule with gonocoxae apically rounded (Fig. 80A, B)61
_	Genital capsule with clearly produced gonocoxal teeth (Fig. 80C–G) 62

¹⁸ These two species are very difficult to separate in the male sex. Association with females should be made, as well as taking account of their known distributions.

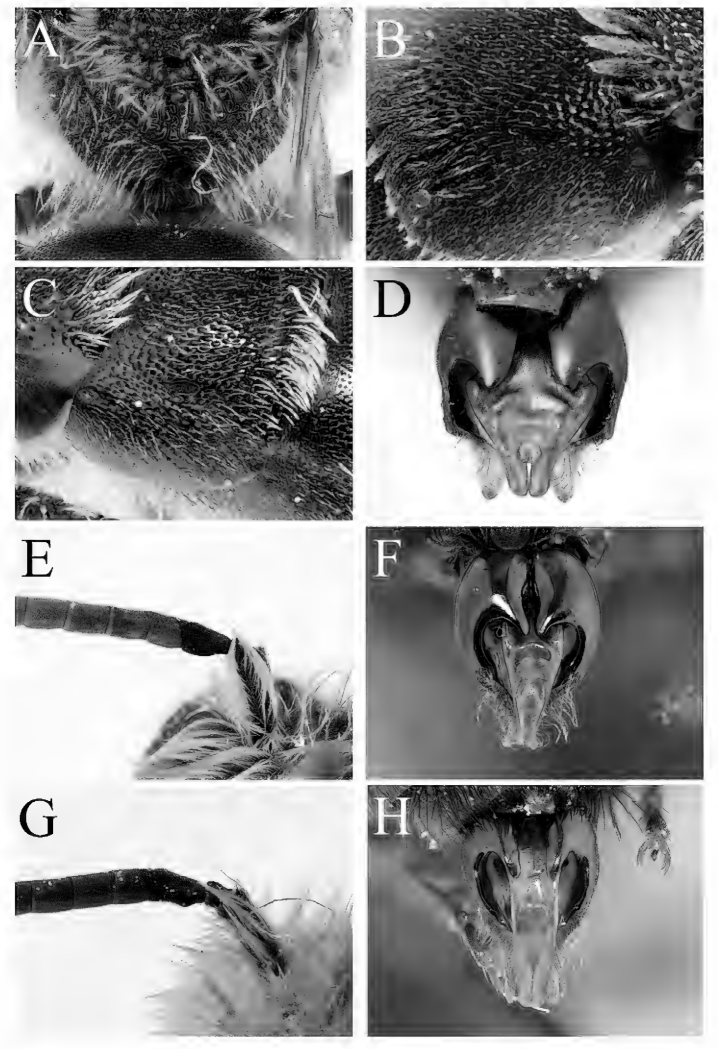


Figure 79. Andrena (Brachyandrena) colletiformis Morawitz, 1873, male **A** propodeum, dorsal view **B** T2, profile view; Andrena (Brachyandrena) miegiella Dours, 1873, male **C** T2, profile view; Andrena (Suandrena) gades Wood & Ortiz-Sánchez, 2022, male **D** genital capsule, dorsal view; Andrena (Suandrena) cyanomicans Pérez, 1895, male **E** antennae, frontal view **F** genital capsule, dorsal view; Andrena (Suandrena) suerinensis Friese, 1884, male **G** antennae, frontal view **H** genital capsule, dorsal view.

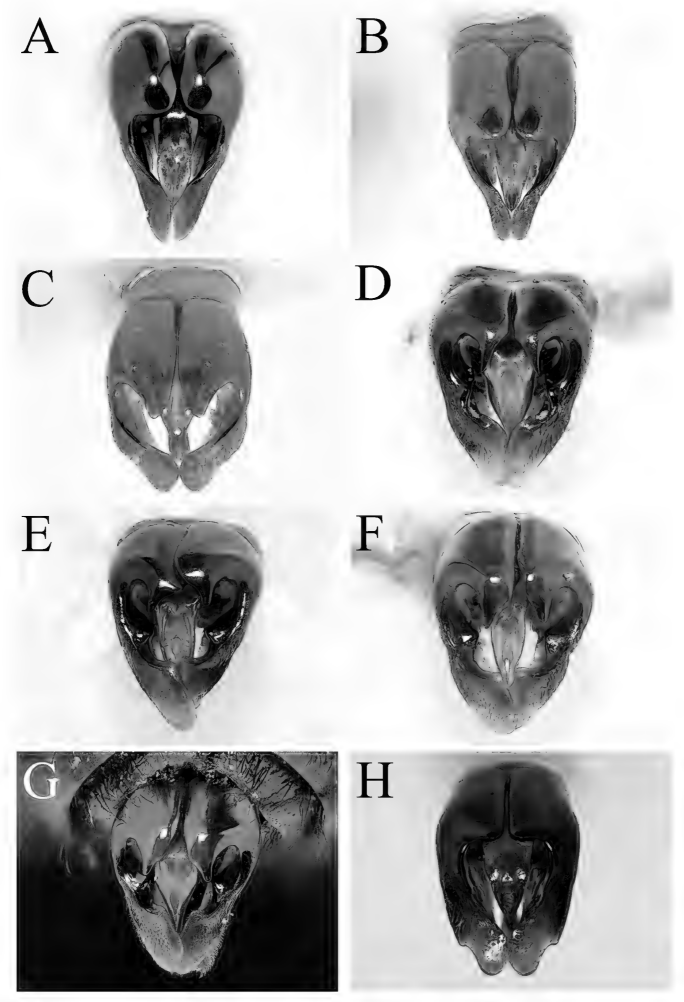


Figure 80. Andrena (Plastandrena) agilissima (Scopoli, 1770), male A genital capsule, dorsal view; Andrena (Plastandrena) afrensis Warncke, 1967, male B genital capsule, dorsal view; Andrena (Plastandrena) asperrima Pérez, 1895, male C genital capsule, dorsal view; Andrena (Plastandrena) tibialis (Kirby, 1802), male D genital capsule, dorsal view; Andrena (Plastandrena) bimaculata (Kirby, 1802), male E genital capsule, dorsal view; Andrena (Plastandrena) pilipes Fabricius, 1781, male F genital capsule, dorsal view; Andrena (Plastandrena) nigrospina Thomson, 1872, male G genital capsule, dorsal view; Andrena (Melandrena) flavipes Panzer, 1799, male H genital capsule, dorsal view.

61	Terga with metallic blue reflections. Genital capsule with penis valves medially with and oval hyaline expansion, this breaking the profile of the penis
	valves themselves (Fig. 80A). Gonostyli with inner margin raised
	agilissima (Scopoli)
_	Terga dark, without metallic blue reflections. Genital capsule penis valves
	uniformly converging apically (Fig. 80B). Gonostyli with inner margin not
	raised
62	Terga usually extremely densely punctate, punctures almost confluent. Ter-
	ga with weak metallic blue hints. Genital capsule, see Fig. 80C, see also
	Fig. 25D
_	Terga never so densely punctate, punctures never confluent, always sepa-
	rated by at least 0.5 puncture diameters. Terga never with metallic blue re-
	flections. Genital capsule otherwise
63	Mesosoma with brown pubescence dorsally and laterally, at most with oc-
03	casional black hairs
	Mesosoma with grey and black pubescence, always with at least some black
_	hairs laterally on the mesepisternum65
64	•
04	Genital capsule with penis valves basally broad (Fig. 80D). Nominally uni-
	voltine, flying only in the spring tibialis (Kirby)
_	Genital capsule with penis valves basally narrow (Fig. 80E). Bivoltine, flying
<i>(</i> =	in the spring and the summer bimaculata (Kirby)
65	Penis valves basally narrow, apical spatulate parts of gonostyli comparative-
	ly narrow (Fig. 80F). Bivoltine (typically March–April and July–August).
	Common and widely distributed throughout Iberia pilipes Fabricius
_	Penis valves basally broad, apical spatulate parts of gonostyli comparative-
	ly broad (Fig. 80G). Univoltine (typically May-June). Rare, restricted to
	mountainous parts of Iberia; known from the Pyrenees, the Sistema Cen-
	tral, Serra do Gerês, Sistema Ibérico, and Sierra de Cazorla
66 (51)	Genital capsule distinctive, with more or less 90° emargination in the outer
	parts of the gonostyli (Fig. 80H). The most common Andrena species in
	Iberia
_	Genital capsule otherwise67
67	Process of the labrum thickened and expanded, anteriorly projecting be-
	yond the fore margin of the clypeus. Pronotum with strong humeral angle.
	Active in the summer (mid-June to September) (subgenus Cnemidandrena
	and remaining Margandrena)
_	Without this combination of characters71
68	T2-3 extensively red-marked. Clypeus with apical corners produced into
	distinct anteriorly projecting knobs (c.f. Fig. 76A). Restricted to north-east-
	ern Spain in Mediterranean habitats
_	Terga dark, without red markings. Clypeus without apically projecting
	knobs

69	Outer surface of the galea smooth and shiny. Associated with Ericaceae, found in Atlantic habitats across northern, central, and western Iberia
_	Outer surface of the galea shagreened and dull
70	Gena posteriorly produced into a strong winged carina. Scutum with abun-
	dant black hairs. Associated with Asteraceae, restricted to northern Spain
	denticulata (Kirby)
_	Gena posteriorly rounded. Scutum at most with occasional black hairs, usu-
	ally entirely brown-haired. Polylectic, though often found on Asteraceae.
	Restricted to areas surrounding the Pyrenees with isolated populations in
	high mountains in southern Spain (particularly the Sierra Nevada)
71 (67)	Fore margin of clypeus upturned AND gena conspicuously broadened, wid-
, 1 (0,)	er than the width of the compound eye AND pronotum with a conspicuous
	humeral angle (remaining <i>Notandrena</i>)
_	Without this exact combination of characters. (Note, male members of the
	Ovandrena are very similar, but the gena is as broad as the width of the com-
	pound eye, not broader. They can be found at couplet 168; Andrena blanda
	is also similar, but the gena is only slightly wider than the width of the
	compound eye. It should be diagnosed by its genital capsule, go to couplet
	184)
72	Mandibles long, sickle-shaped, strongly crossing apically. A3 long, exceed-
, –	ing length of A4+5. Clypeus covered with dense plumose hairs than obscure
	the underlying surface in fresh specimens. Clypeus sometimes with weak
	metallic reflections
_	Mandibles normal, not sickle-shaped, not strongly crossing apically. A3 short,
	slightly shorter than A4+5. Clypeus without plumose hairs that obscure the
	underlying surface. Clypeus dark, without metallic reflections75
73	Terga uniformly dark, at most with marginal areas lightened dark brown.
	A3 comparatively short, only moderately exceeding length of A4+5. Discs
	of T2-4 clearly and regularly punctate, punctures separated by 2 puncture
	diameters, underlying surface shagreened and weakly shiny
_	Terga red-marked, with at least marginal areas and lateral parts of T2-3
	lightened red. A3 comparatively long, almost equalling or equalling length
	of A4+5+6. Terga shagreened, obscurely punctate, punctures disappearing
	into background sculpture
74	Clypeus polished and shiny, with scattered punctures. Mesonotum with
	shiny areas medially. A3 equalling A4+5+6 binominata Smith
_	Clypeus uniformly shagreened and dull. Scutum uniformly shagreened and
	dull. A3 slightly shorter than A4+5+6leucophaea Lepeletier
75	Terga with marginal areas of T2-4 puncture-free. A3 shorter than A4+5,
	never twice as long as A4. Basitarsi usually dark, sometimes apically lightened

	orange. Restricted to cooler parts of northern and western Iberia; univolt-
	ine, flying on July-August
_	Terga with punctures of discs of T2-4 extending onto marginal areas. A3 al-
	most as long as A4+5, typically twice as long as A4. Basitarsi usually entirely
	lightened orange. Widespread across Iberia; bivoltine, flying in April-May
	and July-August
76 (71)	Small species, body length of 5-7 mm, terga always dark, legs always
	dark (remaining Aciandrena, Aenandrena partim, Avandrena, Cordan-
	drena partim, Cryptandrena partim, Graecandrena, and almost all Mi-
	crandrena)77
_	Larger species, at least 8 mm in length. Terga lightened red in some species,
	either on the discs or the margins. Hind tarsi or basitarsi lightened orange
	in some species
77	All tergal discs uniformly punctate, punctures separated by 0.5 puncture
	diameters, underlying surface weakly shiny. Genital capsule elongate, with
	weakly pronounced gonocoxal teeth, gonostyli apically produced into trian-
	gular wedges (Fig. 77B) ventricosa Dours (partim, dark form)
_	Genital capsule otherwise; tergal punctation variable
78	A3 much shorter than A4, at most ½ the length (Fig. 81A, C). Terga often
	with weak metallic bronzy reflections (Fig. 81D)
_	A3 as long as A4 or longer. Terga without metallic reflections
79	A3 extremely short relative to A4, at most $\frac{1}{5}$ the length (Fig. 81A). Terga
	dark. Genital capsule distinctive, with extremely narrow parallel-sided
	penis valves (Fig. 81B). Rare, known from eastern Spain (Jáen, Soria,
	Teruel)
_	A3 not so short, around ½ the length of A4 (Fig. 81C). Terga with metallic
	bronzy reflections. Genital capsule otherwise80
80	T3-4 laterally almost impunctate, any obscure punctures disappear into the
	background sculpture
_	T3-4 laterally with abundant clear punctures, these distinct against the
	background structure (Fig. 81D)
81	A3 long, equalling or exceeding length of A4+5. Face with abundant dark
	pubescence, sometimes intermixed with white hairs. Propodeal triangle nar-
	row, with fine granular shagreenation over the majority of its area, some-
	times with short, fine, and raised rugae basally (Fig. 82A). Scutum sha-
	greened, dull to weakly shiny, with obscure scattered punctures. Metasoma
	with obscure and scattered hair-bearing punctures, punctures separated by
	3–4 puncture diameters. Active in the early spring (March-April), associated
	with <i>Erodium</i> (Geraniaceae; subgenus <i>Avandrena</i>)
_	Without this combination of characters84
82	Genital capsule relatively compact, gonostyli apically flattened, more or less
	triangular shovel-like, more or less as broad as long (Fig. 82B). Tergal discs

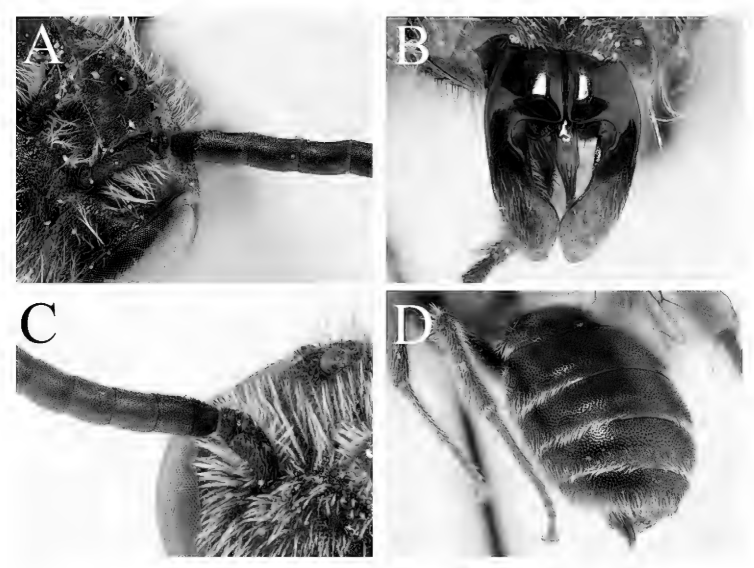


Figure 81. Andrena (Cordandrena) vaulogeri Pérez, 1895, male **A** antennae, frontal view **B** genital capsule, dorsal view; Andrena (Aenandrena) hedikae Jäger, 1934, male **C** antennae, frontal view **D** terga, posterolateral view.

¹⁹ Because of taxonomic complexity and variation in the shape of the genital capsule within *A. avara* (e.g. the capsule pictured here in Fig. 82D is *A. avara liturata* Warncke, 1975 that has apically truncate gonocoxal teeth), it is best to use only the form of S8 to separate *A. panurgina* from the *A. avara* complex.

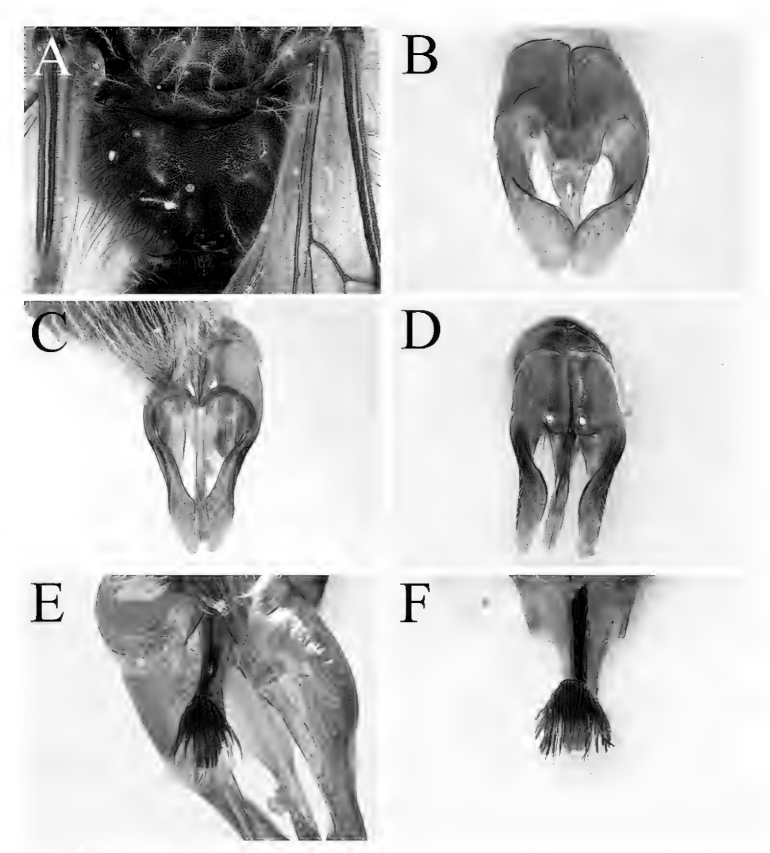


Figure 82. Andrena (Avandrena) panurgina De Steffani, 1889, male **A** propodeal triangle **C** genital capsule, dorsal view **E** S8, ventral view; Andrena (Avandrena) melacana Warncke, 1967, male **B** genital capsule, dorsal view; Andrena (Avandrena) avara liturata Warncke, 1975, male **D** genital capsule, dorsal view **F** S8, ventral view.

85	Clypeus with clear longitudinal striations covering the entire surface. Restricted
	to sandy and usually coastal habitats in southern Iberia orana Warncke
_	Clypeus without striations, or with at most obscure striations at the base of
	the clypeus, never covering the entire surface
86	Clypeus weakly but distinctly domed, basally densely punctate, punctures
	separated by 0.5 puncture diameters, punctures becoming sparse apically,
	here separated by 2-3 puncture diameters; underlying surface smooth and
	shiny (Fig. 83A). Fore margin of the clypeus forming a very wide and gen-
	tle emargination, process of the labrum narrow, slightly longer than wide.
	Genital capsule, see Fig. 83B pandosa trigona Warncke
_	Without this combination of characters, clypeus usually flattened and/or
	shagreened and dull, or genital capsule otherwise87
87	Terga strongly and clearly punctate, tergal margins strongly depressed, with
	punctures continuing onto marginal areas; underlying surface finely sha-
	greened and weakly shiny. Very small species, not exceeding 6 mm
	fulica Warncke (partim, dark form with black clypeus)
_	Terga either impunctate (Fig. 52E) or obscurely punctate, without clear
	punctures extending onto the tergal margins88
88	Clypeus flattened, shagreened in basal half, finely shagreened in apical half,
	the two half therefore contrasting; surface with obscure and scattered punc-
	tures. Genital capsule elongate, with long and sharply pointed penis valves
	(Fig. 83C)verticalis Pérez
_	Clypeus flattened or domed, but without this combination of characters.
	Genital capsule otherwise89
89	Genital capsule unusual, gonocoxae with inner margins forming obtuse
	angle, without gonocoxal teeth, gonostyli flattened with square truncate
	apexes (Fig. 83D, E)90
_	Genital capsule otherwise91
90	Genital capsule with gonostyli comparatively narrow, apically narrower than
	the basal width of the penis valves; penis valves occupying comparatively lit-
	tle of the space between the gonostyli (Fig. 83D). Restricted to the extreme
	north-east of Spainimpunctata Pérez
_	Genital capsule with gonostyli comparatively broad, apically as broad as the ba-
	sal width of the penis valves; penis valves occupying majority of space between
	the gonostyli (Fig. 83E). Restricted to central, southern, and eastern Spain, not
	known from the north-east
91	Tergal margins extensively lightened hyaline-yellow, T2-4 with dense and
	thick complete apical hair bands that occupy and slightly exceed the entire
	length of the margin, obscuring the underlying surface in fresh individu-
	als. Small species, not exceeding 6 mm. Rare, known only from central
	Spain
_	Tergal margins with weaker apical hair bands, tergal margins not so exten-
	sively lightened yellow. If in doubt, species exceeding 6 mm in length, or
	with clypeus flattened92

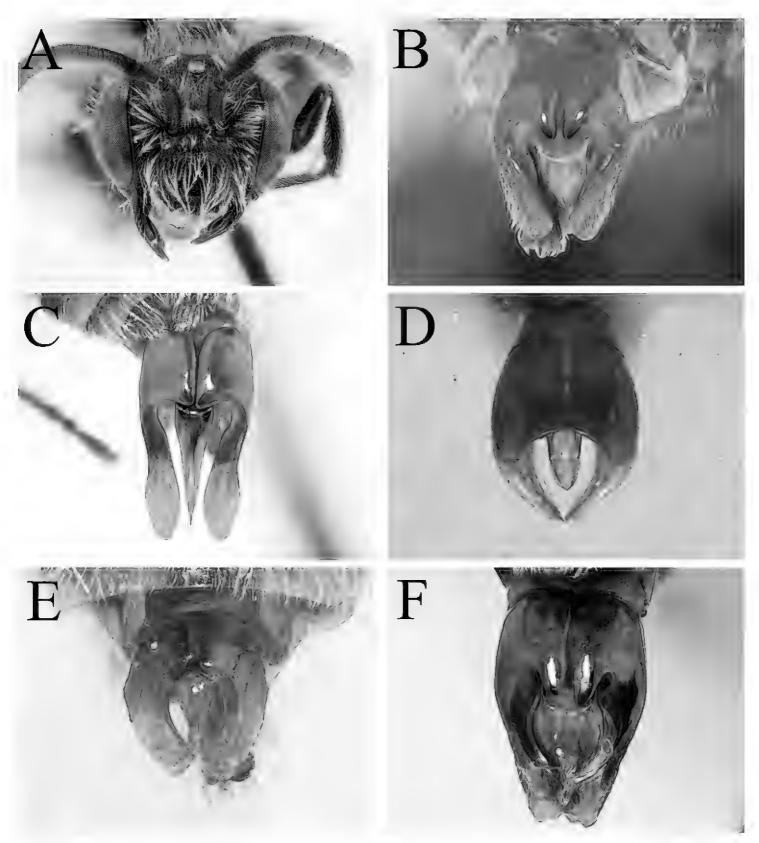


Figure 83. Andrena (Micrandrena) pandosa trigona Warncke, 1975, male **A** face, frontal view **B** genital capsule, dorsal view; Andrena (Graecandrena) verticalis Pérez, 1895, male **C** genital capsule, dorsal view; Andrena (Graecandrena) impunctata Pérez, 1895, male **D** genital capsule, dorsal view; Andrena (Graecandrena) nebularia Warncke, 1975, male **E** genital capsule, dorsal view; Andrena (Micrandrena) longibarbis Pérez, 1895, male **F** genital capsule, dorsal view.

93	A3 shorter than A4+5. Clypeus flattened, with slight metallic green- purple hints medially. Terga essentially impunctate. Widespread across Iberia
_	A3 slightly longer than A4+5. Clypeus flattened, almost with slight longitudinal concavity medially, uniformly black. Terga obscurely punctate. Central and eastern Spain only
94	Gena broad, clearly broader than the width of a compound eye (Fig. 52B, C). Scutum with obscure and shallow but large, hair-bearing punctures, punctures separated by 2–3 puncture diameters. Pronotum with strong humeral angle. Widespread across western and southern Iberia
_	Gena normal, equalling the width of a compound eye. Scutum extremely obscurely punctate, punctures disappearing into background structure. Pronotum rounded. Presence and distribution in Iberia unclearabjecta Pérez
95	Genital capsule distinctive, with strongly reflexed gonostyli (Fig. 84A) djelfensis Pérez
_	Genital capsule otherwise96
96	Genital capsule with penis valves clearly and strongly broadened basally, clearly bulbous in appearance (Fig. 84B–D)
_	Genital capsule with penis valves not strongly broadened basally100
97	Marginal area of T1 wide, strongly thickened, impunctate. Scutum with scattered punctures, punctures separated by 1–3 puncture diameters. Genital capsule with gonocoxal teeth weakly produced (Fig. 84B) Restricted to mountainous areas in northern Spain, associated with <i>Potentilla</i> (Rosace-
	ae)
_	Marginal area of T1 not strongly thickened, flat or slightly depressed. Genital capsule with gonocoxal teeth truncate (Fig. 84C, D). Scutum densely punctate, punctures separated by 0.5–1 puncture diameter98
98	Clypeus flattened, with slight longitudinal impression medially. Inner mar-
	gin of the gonostyli upturned, diverging dorsally from outer margin, forming a slight but visible 'kink' (Fig. 84C). Associated with <i>Ornithogalum</i> (As-
	paragaceae)
_	Clypeus less strongly flattened. Inner margin of the gonostyli parallel with the outer margin, without a visible 'kink' (Fig. 84D). Associated with a
	wider variety of flowering plants99
99	Restricted to areas close to the Pyrenees. A4 slightly longer than A6. Broadened penis valves comparatively longer strokenst strokenst Stöckhert
_	Found in mountainous areas across southern and central Spain and northern Portugal. A4 as long as A6. Broadened penis valves comparatively shorter ²⁰

²⁰ These two species are very difficult to separate morphologically; geographic context should be used.

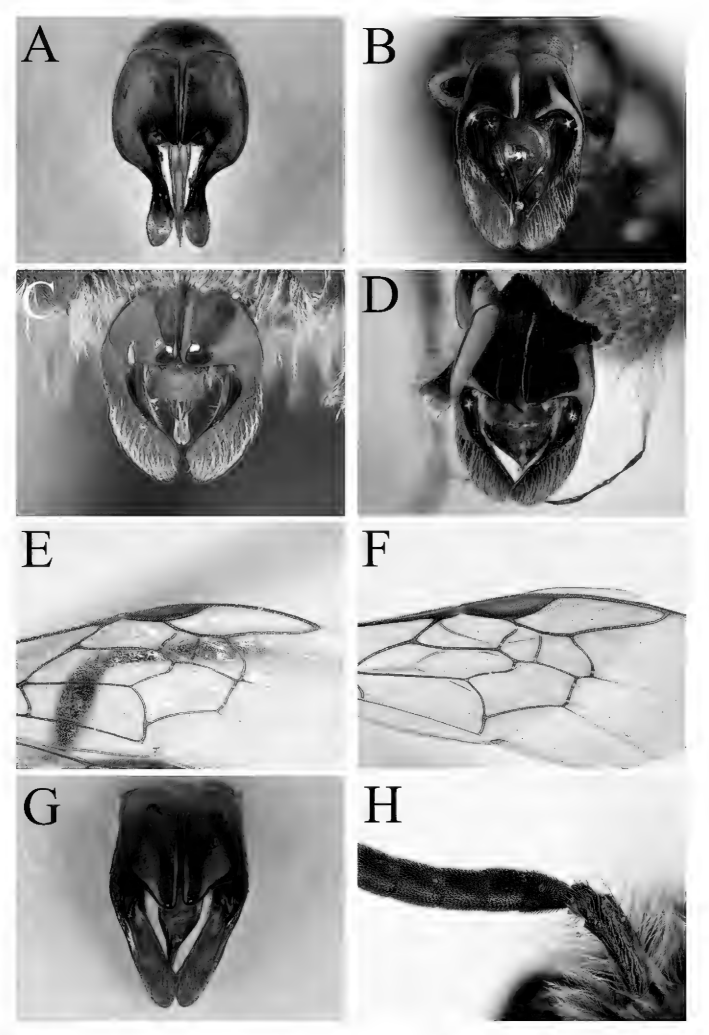


Figure 84. Andrena (Micrandrena) djelfensis Pérez, 1895, male **A** genital capsule, dorsal view; Andrena (Micrandrena) falsifica Perkins, 1915, male **B** genital capsule, dorsal view; Andrena (Micrandrena) saxonica Stöckhert, 1935, male **C** genital capsule, dorsal view; Andrena (Micrandrena) icterina Warncke, 1974, male **D** genital capsule, dorsal view; Andrena (Micrandrena) nana (Kirby, 1802), male **E** forewing, dorsal view **F** Andrena (Micrandrena) fabrella Pérez, 1903, male **F** forewing, dorsal view **G** genital capsule, dorsal view; Andrena (Micrandrena) semilaevis Pérez, 1903, male **H** antennae, frontal view.

100	Genital capsule with outer margins of the gonostyli concave. Clypeus
	strongly shagreened and dull simontornyella Noskiewicz
_	Genital capsule with outer margins of the gonostyli essentially parallel with
	the inner margin101
101	Clypeus flattened, basally with obscure but distinct longitudinal striations,
	these covering the basal quarter of the clypeus and are channelled laterally
	along its margins102
_	Clypeus typically domed, without longitudinal striations basally103
102	Facial pubescence predominantly black. Scutum shagreened with faint and
	weak greasy metallic reflections. Tergal discs shagreened, weakly shiny. Tergal
	margins distinctly depressed. Widespread throughout Iberia nitidula Pérez
_	Facial pubescence predominantly pale, with scattered black hairs laterally.
	Scutum more strongly shagreened, matt. Tergal discs with strong microre-
	ticulation, dull. Tergal margins weakly depressed. Restricted to the extreme
	north-east of Spain
103	Genital capsule elongate with penis valves converging apically to form a
	tapering point (Fig. 20A, B). Propodeum with propodeal triangle broad,
	laterally poorly defined with lateral carinae becoming shorter and weaker
	posteriorly104
_	Genital capsule and propodeal triangle otherwise105
104	Genital capsule with the inner margins of the gonostyli only weakly pro-
	duced towards the penis valves; apical spatulate part of the gonostyli com-
	paratively elongate (Fig. 20A). Tergal discs typically more shagreened and
	dull, punctures disappearing somewhat into underlying sculpture
_	Genital capsule with the inner margins of the gonostyli strongly produced
	towards the penis valves; apical spatulate part of the gonostyli comparatively
	short (Fig. 20B). Tergal discs typically smooth and shiny, punctures clearly
	visible
105	Tergal discs, at least on T1, smooth and shiny, at most with very fine sha-
	greenation106
_	Tergal discs shagreened or microreticulate, not strongly shiny, most dull and
_	matt113
106	Forewing with first submarginal cross vein separated from the stigma by
	three times its own width (Fig. 84E)107
_	Forewing with first submarginal cross vein separated from the stigma by
	more or less its own width (Fig. 84F)108
107	Clypeus medially shagreened. Stigma dark brown. Sterna shagreened with
	the exception of the marginal zones. Common and widespread across
	Iberianana (Kirby)
_	Clypeus medially shiny. Stigma bright, yellowish to reddish. Sternal discs
	shiny. Very rare, restricted to the Pyrenees
108	Scutum densely shagreened, with large shallow 'crater-punctures'. In fresh
	specimens, T1-5 with broad white apical hair bands that exceed the length

	of the marginal areas. S2–5 with distinct white apical hair bands. Restricted
	to mountainous areas of central and northern Spain
	Scutum shiny or obscurely shiny, never densely shagreened, with normal
_	punctures without raised rims. Terga and sterna without clear and dense
	complete hair bands109
109	Dorsal part of gena completely smooth and shiny, without any microscu-
10)	lpture. Discs of T2–4 very densely punctate, punctures separated by 0.5
	puncture diameters. Restricted to dry and steppic habitats in central Iberia
	including northern Portugal
_	Gena uniformly shagreened, never with extensive smooth and shiny areas.
	Discs of T2–4 less densely punctate, punctures separated by at least 1 punc-
	ture diameter
110	Antennae extensively lightened orange ventrally. T1 with very fine and scat-
	tered punctation. Rare, restricted to northern Spain, univoltine (August-
	September) nanula Nylander
_	Antennae dark ventrally. T1 with clearer and denser punctation, punctures
	separated by 1-2 puncture diameters
111	Terga with punctation clearly extending onto the depressed marginal ar-
	eas. Genital capsule, see Fig. 84G. Associated with Cistaceae
_	Terga with marginal areas essentially impunctate, at most with occasional
	puncture. Genital capsule otherwise112
112	Marginal areas of T2–4 strongly depressed, mirror-smooth, without a single
	puncture. A4 as long as broad, equalling A3 in length. Found in the Sistema
	Central to west and north-western Iberia. Associated with Sedum (Crassu-
	laceae)
_	Marginal areas of T2–4 comparatively weakly depressed, smooth and shiny,
	but with occasional punctures. A4 subquadrate, broader than long, shorter
112	than A3 in length. Widespread in Iberia
113	A4 quadrate, as wide as long (Fig. 84H). Tergal margins strongly depressed and polished, mirror-smooth, impunctate. Restricted to the Pyrenees and
	Cantabrian Mountainssemilaevis Pérez
	A4 subquadrate, broader than long. Tergal margins depressed or not 114
- 114	Propodeal triangle laterally poorly defined, without clear carinae, internal
111	surface with raised rugosity covering only central part in the form of a tri-
	angle, lateral parts therefore with granular shagreenation (Fig. 85A). Facial
	hair predominantly black. Clypeus usually smooth and shiny in its apical
	half. Terga essentially impunctate, regularly shagreened and dull, tergal
	margins weakly depressed. Univoltine (February–April)
	tenuistriata Pérez
_	Without this exact combination of characters115

115	Scutum strongly shagreened and dull, very obscurely punctate, punctures
	separated by 2-4 puncture diameters, underlying surface matt; punctures
	often disappear into background sculpture116
_	Scutum less strongly shagreened to shiny, with at least some punctures
	clearly visible against the background sculpture; punctures separated by 1-2
	puncture diameters117
116	Tergal discs finely shagreened and weakly shiny, with scattered punctures
	that are nevertheless distinct against the background sculpture, punctures
	separated by 1-3 puncture diameters. Tergal margins with finer shagreen,
	impunctate, thus contrasting the tergal discs. Restricted to temperate areas
	close to the Pyrenees
_	Tergal discs strongly shagreened and dull, with extremely obscure punctures
	that are hard to distinguish against the background sculpture, separated by
	2–4 puncture diameters. Tergal margins with equally strong shagreen, not
	strongly contrasting the tergal discs. Iberian distribution unclear, but prob-
	ably restricted to southern Spain (Cádiz, Málaga)tiaretta Warncke
117	T1–3 with discs strongly and coarsely punctate, punctures clearly visible
	against the background sculpture (Fig. 85B). Tergal margins strongly de-
	pressed, finely shagreened, more or less shiny, thus strongly contrasting the
	tergal discs. Restricted to central and northern Spain, typically at altitude in
	mountainous areas
_	T1-3 without strong and coarse punctures, at most with occasional punc-
	tures that typically disappear into the background sculpture. Tergal margins
	depressed or not, but not noticeably more finely sculptured or more brightly
	shiny than the tergal discs
118	T2-4 with tergal margins strongly depressed119
_	T2-4 with tergal margins at most weakly depressed120
119	Scutum with scattered punctures, punctures separated by 1-3 puncture diam-
	eters. Mesopleuron finely microreticulate, punctate, punctures separated by >1
	puncture diameter. Throughout Iberia, common ²¹ spreta Pérez
_	Scutum with denser punctures, punctures separated by 1-2 puncture diameters.
	Mesopleuron more coarsely microreticulate, punctate, punctures separated by
	<1 puncture diameter. Exact Iberian distribution unclear, but currently known
	only from eastern Spain, typically in mountainous areas ²¹ pauxilla Stöckhert
120	Scutum densely punctate, punctures typically separated by 1 puncture diam-
	eter. Scutum and scutellum shagreened and dull (1st generation) to weakly
	shiny (2 nd generation). Genital capsule, see Fig. 85C minutula (Kirby)
_	Scutum more sparsely and irregularly punctate, punctures separated by 1-3
	puncture diameters. Scutum and scutellum finely shagreened and weakly

²¹ The differentiation between these two species in Iberia in the male sex is not completely clear due to lack of available Iberian male material. Association with females should be made.

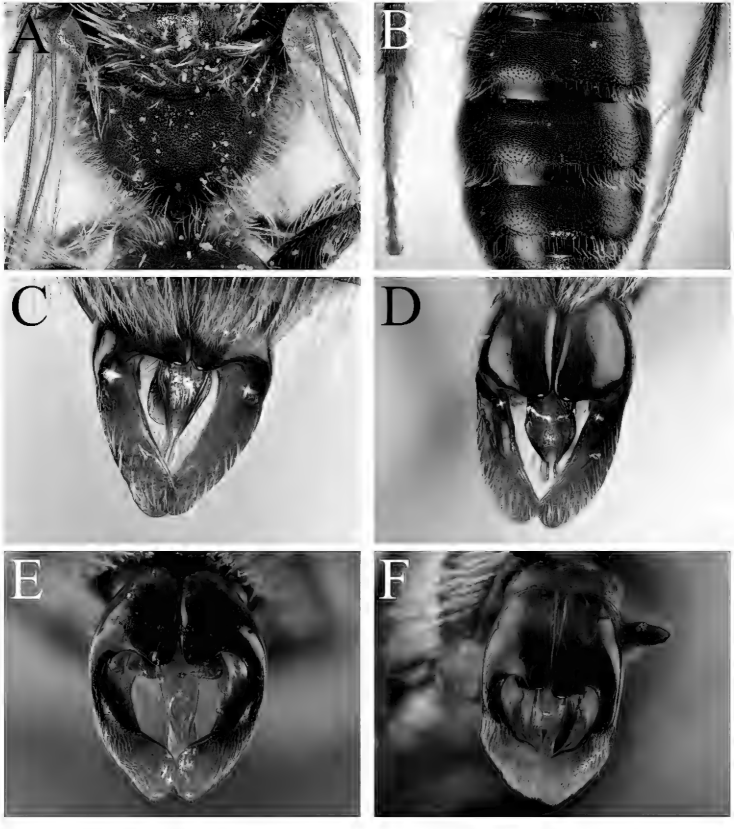


Figure 85. Andrena (Micrandrena) tenuistriata Pérez, 1895, male **A** propodeum, dorsal view; Andrena (Micrandrena) anthrisci Blüthgen, 1925, male **B** terga, dorsal view; Andrena (Micrandrena) minutula (Kirby, 1802), male **C** genital capsule, dorsal view; Andrena (Micrandrena) minutuloides Perkins, 1914, male **D** genital capsule, dorsal view; Andrena (Pruinosandrena) nilotica Warncke, 1975, male **E** genital capsule, dorsal view. **F** Andrena (Pruinosandrena) pruinosa Erichson, 1835, male **F** genital capsule, dorsal view.

122	Dorsolateral parts of the propodeum impunctate, with dense network of raised rugosity
_	Dorsolateral parts of the propodeum regularly punctate, without a dense network of raised rugosity
123	Face and mesosoma with a mixture of black and white pubescence. T2–4 laterally with widely separated patches of dense white pubescence on their apical margins that obscure the underlying surface. Terminal fringe black lateralis Morawitz
_	Face and mesosoma with light brown pubescence. T2–4 laterally without patches of white pubescence, at most with obscure whitish hair bands. Terminal fringe light brown
124	A3 exceeding A4+5 in length
_	A3 not exceeding A4+5
125	Genital capsule without clear kink in the inner margins of the gonostyli (Fig. 85E). Scutum less densely punctate, punctures separated by 0.5 puncture diameters, with shiny interspaces, predominantly shiny
_	Genital capsule with clear kink in the inner margins of the gonostyli (Fig. 85F). Scutum more densely punctate, punctures almost confluent,
	predominantly dull
126 (121)	Clypeus flattened over majority of its surface (subgenus <i>Taeniandrena</i>) ²² 127
_	Clypeus not noticeably flattened
127	Genital capsule with pronounced gonocoxal teeth, these clearly projecting anteriorly for a distance greater than the diameter of an antenna (Fig. 86A, B)
_	Genital capsule with at most weakly projecting gonocoxal teeth (Fig. 86C–F)
128	Penis valves more or less parallel-sided (Fig. 86A). Gonostyli basally (before flattened apical parts) with dense network of longitudinal striations on their dorsal surface. Known only from central Portugal (Castelo Branco) and south-western Spain (Huelva, Cádiz). Flying early in the season (March-April)
_	Penis valves very broad basally, occupying almost entire space between gonostyli (Fig. 86B). Gonostyli without longitudinal striations on their dorsal surface. Restricted to northern Spain. Flying slightly later (April–June) <i>lathyri</i> Alfken

²² Males of this subgenus are highly challenging and in some cases impossible to identify. There are several outstanding taxonomic issues in this subgenus that require in-depth genetic study to resolve, and additional probably undescribed cryptic species are present. Do not expect to identify this group without consulting confidently determined reference material, and ideally barcoded specimens. Minimal characters are given here because of the ongoing lack of taxonomic clarity in this subgenus.

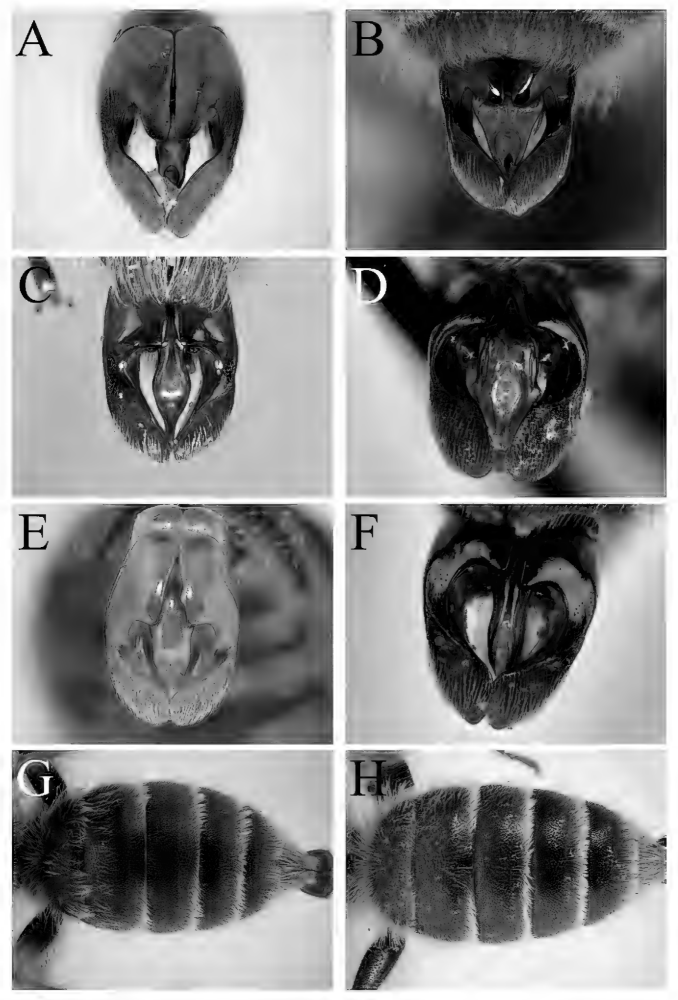


Figure 86. Andrena (Taeniandrena) lusitania Wood & Ortiz-Sánchez, 2022, male A genital capsule, dorsal view; Andrena (Taeniandrena) lathyri Alfken, 1900, male B genital capsule, dorsal view; Andrena (Taeniandrena) gredana Warncke, 1975, male C genital capsule, dorsal view; Andrena (Taeniandrena) intermedia Thomson, 1870 aggregate, male D genital capsule, dorsal view; Andrena (Taeniandrena) levante Wood & Praz, 2021, male E genital capsule, dorsal view; Andrena (Taeniandrena) contracta Wood, 2022, male F genital capsule, dorsal view; Andrena (Taeniandrena) wilkella (Kirby, 1802), male G terga, dorsal view; Andrena (Taeniandrena) benoisti Wood & Praz, 2021, male H terga, dorsal view.

129	Genital capsule with penis valves narrow basally, broad medially, and nar-
	rowing apically (Fig. 86C). Restricted to the Sistema Central and northern
	and western Iberiagredana Warncke
_	Genital capsule different130
130	Penis valves broadened and gonocoxae with their inner margins diverging
	from their base, gonocoxae therefore forming a broad triangular opening
	basally (Fig. 86D, E)
_	Penis valves not broadened or gonocoxae with their inner margins more or
	less parallel, sometimes slightly diverging apically132
131	Penis valves strongly broadened, occupying the majority of the space between
	the gonostyli (Fig. 86D). Gonostyli with strongly raised inner margin, this
	forming a distinct obtuse angle medially. Found across mountainous areas in
	eastern Spain
	likely represents a complex of an unknown number of species, potentially all
	of which are undescribed; the true <i>intermedia</i> may be absent from Spain)
_	Penis valves not so strongly broadened, occupying ½ the space between
	the gonostyli (Fig. 86E). Gonostyli with weakly raised inner margin,
	not presenting a distinct angle. Restricted to southern and south-eastern
	Spain (Almería, Granada, Málaga, Murcia, Valencia). Univoltine (April-
	June)
132	Genital capsule with weakly produced gonocoxal teeth (Fig. 86F). Penis
	valves very narrow, more or less parallel-sided. Restricted to the alpine zone
	of the Sierra Nevada (typically above 2000 m)
_	Genital capsule different, species found elsewhere133
133	A3 short relative to A4, approximately 0.6–0.7 times as long134
_	A3 long, more or less as long as A4, typically between 0.8–1.2 times as
	long
134	Clypeus densely covered with snow-white vestiture, in fresh specimens com-
	pletely obscuring the underlying surface, particularly of the apical margin.
	Terga strongly shagreened, punctation obscure, T1 with declivity almost im-
	punctate, any punctures sparse and obscure. Marginal areas of T2–4 almost
	impunctate. Currently known only from the steppe of central Spain (Guada-
	lajara, Salamanca, Segovia). Univoltine (May–June) ovata Schenck
_	Clypeus without such pubescence, the surface of the clypeus is usually clear-
	ly visible through the sparser pubescence. Terga with stronger more distinct
	punctation, including declivity of T1. Marginal areas of T2-4 distinctly
	punctate, at least basally135
135	Terga with weak apical hair bands, widely interrupted on T2–3, complete
	on T4 (Fig. 86G). Smaller, 8–9 mm. Restricted to northern Portugal and
	Spain to the Pyrenees with isolated populations at elevation in the Sistema
	Central and Sistema Ibérico
_	Terga with strong and thick apical hair bands, medially interrupted on T2,
	complete on T3–4 (Fig. 86H). Larger, 10–11 mm. Restricted to the Sis-
	tema Central to central and northern Portugal and north-western Spain

	(Zamora). Not known from the Pyrenees or Cantabrian Mountains
136	Hind tibiae and basitarsi lightened orange. Terga strongly shagreened, usually with punctures obscure and disappearing into underlying sculpture. A3 sometimes a little longer than A4. Larger, 10–12 mm. Univoltine (April–June)russula Lepeletier sensu lato (including the distinct mitochondrial lineage from southern Portugal)
_	Hind legs usually dark. Terga variable, often with abundant and distinct punctures. A3 variable. Smaller, 8–10 mm. Bivoltine (March-August)
137	A3 often shorter than A4 (Fig. 87E). Genital capsule with gonostyli produced into apical points (Fig. 87F). Currently only confirmed from the coast of southern Spain (Málaga). Bivoltine (probably February–April; May–June) ²³
_	Genital capsule with gonostyli apically truncate, not produced into points. 138
138	A4 usually equal or subequal to A3 in length (Fig. 87A). Terga less densely punctate, surface more finely shagreened, weakly shiny. Genital capsule comparatively less elongate, gonostyli broad with external margin usually straight (Fig. 87B). Internal margins of gonocoxae slightly divergent apically. Penis valve slightly broader basally. Throughout Iberia. Bivoltine (typically May–June; July–August)
139 (126)	June–July)
_	Tergal discs never red-marked, at most with tergal margins lightened 144
140	A3 longer than A4
_	A3 much shorter than A4, at most ¼ the length143
141	Head broad, inner margin of compound eyes diverging ventrally (Fig. 41F). Pronotum with strong humeral angle. Clypeus smooth and shiny with scattered punctures. Genital capsule unique, elongate, gonocoxae truncate with inner margin rounded, penis valves basally broad, strongly narrowing medially to become elongate and acutely pointed apically (Fig. 41H)
_	Without this combination of characters

²³ These three taxa are extremely challenging to separate in the male sex. Comparison to confidently determined or preferably barcoded specimens should be made, as well as association with females.

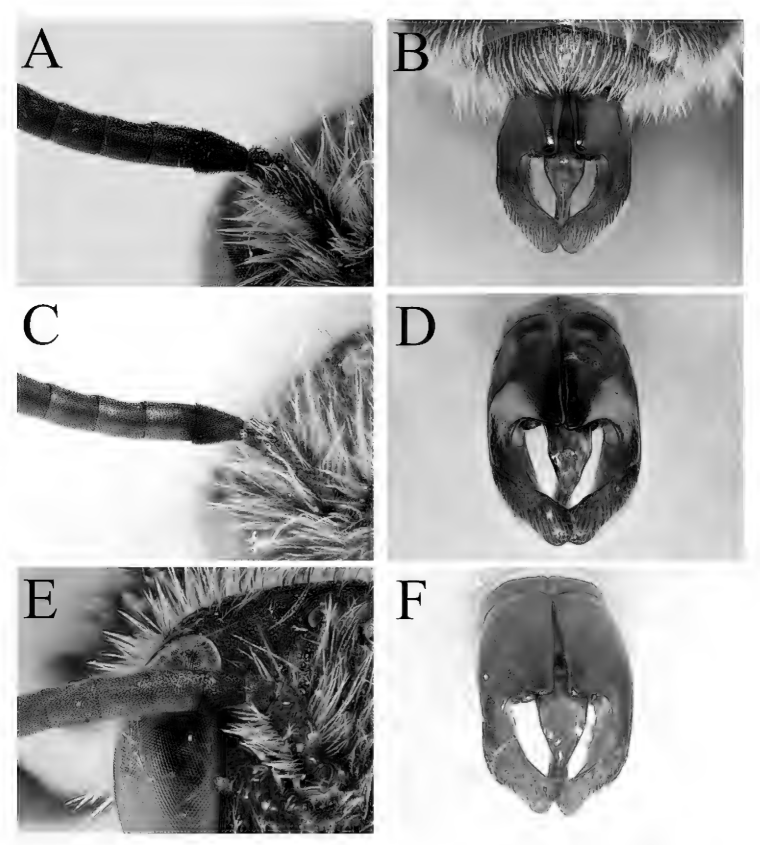


Figure 87. Andrena (Taeniandrena) afzeliella (Kirby, 1802), male **A** antennae, frontal view **B** genital capsule, dorsal view; Andrena (Taeniandrena) ovatula (Kirby, 1802), male **C** antennae, frontal view **D** genital capsule, dorsal view; Andrena (Taeniandrena) poupillieri Dours, 1872, male **E** antennae, frontal view **F** genital capsule, dorsal view.

143	S8 apically truncate, lacking apical emargination. Mandibles always biden-
	tate
_	S8 with clear apical emargination. Mandibles unidentate (1st generation) or
	bidentate (2 nd generation)trimmerana (Kirby) (partim, light form)
144 (139)	Tergal discs with metallic blue reflections
_	Tergal discs dark, without metallic reflections, or at most with greasy greenish
	or bronzy reflections laterally (if greenish reflections present on clypeus, scutum,
	and scutellum, go to couplet 182), never with blue metallic reflections 147
145	Tergal discs with abundant 'crater punctures' with raised rims. Pubescence
	of mesosoma and face light brown, never with abundant black and white
	hairs
_	Tergal discs with simple punctures, without noticeably raised rims. Pubes-
	cence of mesosoma with abundant black and white hairs, on the mesosoma
	forming a striped pattern; white anteriorly and posteriorly, medially with a
	uniform black strip
146	A3 0.6 times as long as A4. Facial hair shorter than the width of a com-
	pound eye. Antennal scape with intermixed black and white hairs. Bivoltine
	(April-May; July-August), restricted to mountains in northern Spain and
	the Pyrenees ²⁴
_	A3 as long or almost as long as A4. Facial hair exceeding the width of a com-
	pound eye. Antennal scape with uniformly white hairs. Univoltine (April–May),
	more widespread across northern Portugal and Spain ²⁴ cineraria (Linnaeus)
147 (144)	Mandibles elongate, sickle-like, strongly crossing apically (Fig. 88A).
` ,	Pronotum with strong humeral angle. Gena often expanded, broader than
	the width of a compound eye148
_	Without this combination of characters; mandibles normal, not elongate
	and strongly crossing apically (Fig. 88B). Pronotum usually without a strong
	humeral angle. Gena usually normal, equalling or shorter than the width of
	a compound eye
148	Genital capsule distinctive, gonostyli long and filiform, penis valves gross-
	ly expanded, occupying almost the entire space between the gonostyli
	(Fig. 88C). S8 elongate and strongly emarginate medially, forming a 'Y'
	shape. Rare, restricted to the Pyreneesbucephala Stephens
_	Genital capsule and S8 otherwise
149	A3 extremely short, at most 1/4 the length of A4
_	A3 longer, never this short. If in doubt, then terga with well-defined and
	narrow light brown apical hair bands153
150	S8 apically truncate, lacking apical emargination. Mandibles always biden-
	tate
_	S8 with clear apical emargination. Mandibles either unidentate or biden-
	tate

²⁴ These two species are challenging to separate morphologically.

151	Mandible unidentate, lacking an inner subapical tooth. Gena usually with a long spine. Flying only in the spring (March-May, depending on eleva-
	tion) trimmerana (Kirby) (partim, dark form, 1 st generation)
	Mandible bidentate, with an inner subapical tooth. Gena usually without a
_	<u> </u>
	spine, sometimes with a very short spine. Flying in the spring or the sum-
150	mer
152	Flying in the spring (usually April to mid-June). Facial pubescence long.
	Rare, restricted to cooler parts of Iberiascotica Perkins
_	Flying in the summer (usually mid-June to July). Facial pubescence short. Widespread throughout Iberia
	trimmerana (Kirby) (partim, dark form, 2 nd generation)
153	Mandible unidentate, without inner subapical tooth. Sometimes there may
	be a hint of a weakly formed or ancient subapical tooth; this state should be
	treated as unidentate. Head often grossly enlarged154
_	Mandible clearly bidentate, with a strong inner subapical tooth. Head not
	normally grossly enlarged156
154	A3 slightly shorter than A4. Clypeus in apical third bulging, surface smooth
	and shiny with scattered punctures, punctures separated by 1-4 punc-
	ture diameters. Hind tibiae and tarsi usually lightened orange. Northern
	and central Spain, in areas with deciduous forest. Associated with Quercus
	(Fagaceae)
_	A3 clearly longer than A4. Clypeus even, without bulging apical part, un-
	derlying surface evenly shagreened and regularly punctate. Hind tibiae and
	tarsi dark
155	Clypeus densely punctate, punctures separated by 0.5–1 puncture diameter.
- , ,	Clypeus apically occasionally with very small pale marking medially. Tergal
	discs with scattered punctures, punctures separated by 2–3 puncture diam-
	eters, underlying surface smooth and shiny. Bivoltine, flying April-May and
	again July-August ²⁵
_	Clypeus with sparser punctures, punctures separated by 1–2 puncture di-
	ameters. Tergal discs densely punctate, punctures separated by 1–2 puncture
	diameters, underlying surface shagreened, weakly shiny. Univoltine, typi-
	cally flying March-June depending on altitude ²⁵ vetula Lepeletier
156	
170	A3 distinctly shorter than A4. Genital capsule with gonocoxal teeth pro-
	nounced but strongly truncate apically, quadrangular (Fig. 88D). Terga with
	well-defined and narrow apical light brown hair bands, these occupying only
	the apical part of the marginal areas (Fig. 88E)leptopyga Pérez
_	A3 at least as long as A4, usually longer. Genital capsule either with less
	strongly pronounced gonocoxal teeth, or with gonocoxal teeth apically
	rounded. Terga with or without apical tergal hair bands157

²⁵ These two species are unrelated, but have converged on a quite similar morphology; more care is required than might be expected based on their divergent female morphology.

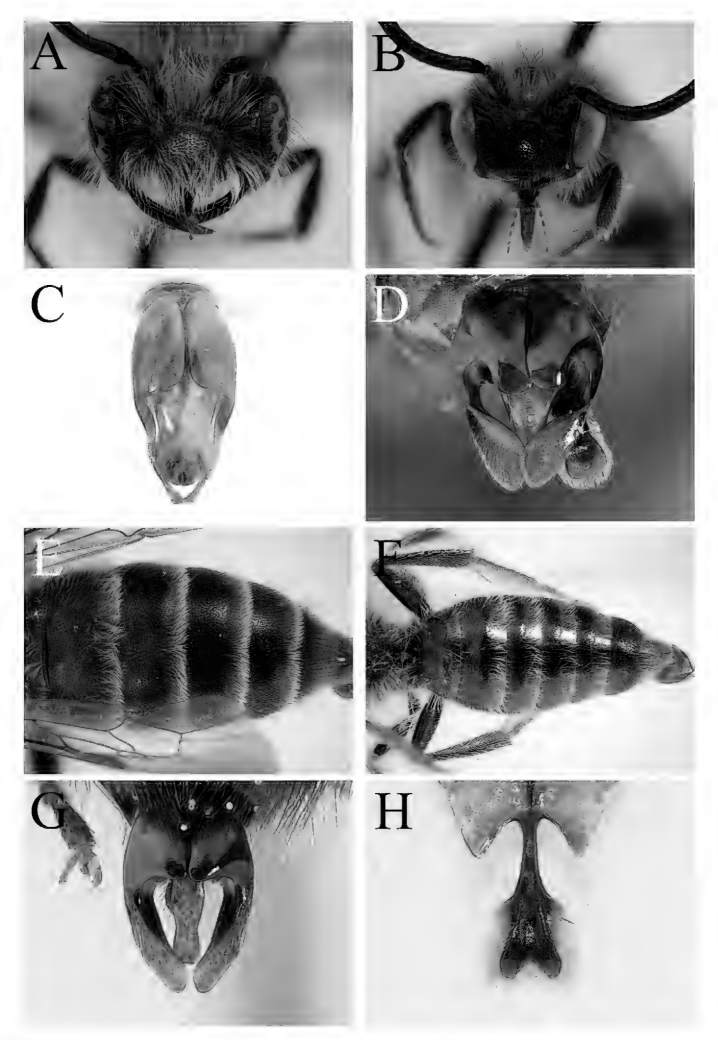


Figure 88. Andrena (Andrena) helvola (Linnaeus, 1758), male **A** head, frontal view; Andrena (Chrysandrena) fulvago (Christ, 1791), male **B** head, frontal view; Andrena (Hoplandrena) bucephala Stephens, 1846, male **C** genital capsule, dorsal view; Andrena (Leucandrena) leptopyga Pérez, 1895, male **D** genital capsule, dorsal view **E** terga, dorsal view; Andrena (Euandrena) angustior (Kirby, 1802), male **F** tergal, dorsal view; Andrena (Euandrena) lavandulae Pérez, 1902, male **G** genital capsule, dorsal view; Andrena (Lepidandrena) paucisquama Noskiewicz, 1924, male **H** S8, dorsal view.

157	Marginal area of T2 broad, occupying ½ the length of the segment, strongly depressed, surface smooth and shiny (Fig. 88F). Found in areas with
	a temperate Atlantic climate across central, north-western, and northern
	Iberia
_	Marginal area of T2 never this broad; at most weakly depressed, surface
	structurally not clearly differentiated from the disc, never smooth and
	shiny
150	Genital capsule with gonocoxae apically rounded, gonocoxal teeth very slightly
158	
	protruding at most (Fig. 88G). Facial pubescence intermixed light brown and
	black, brown medially, black apically and laterally. Usually smaller, 8–9 mm.
	Found in areas with a Mediterranean climate
_	Genital capsule with more strongly projecting gonocoxal teeth, overall
	capsule more robustly built. Facial pubescence either black or pale, never
	mixed. Usually larger, 9–12 mm. Typically not found in areas with a Medi-
	terranean climate, species of temperate deciduous forest (subgenus Andrena
	s. str.)
159	Mandible at its base with a clearly pronounced tooth
_	Mandible at its base with either an angulation or straight165
160	A3 1.8 times longer than A4. Hind tarsi lightened orange. With a patchy
	distribution from northern Portugal and Spain to the Pyrenees, with an
	isolated population in the Sierra de Cazorla (Jáen)
_	A3 at most 1.3 times longer than A4, usually more or less equal in length.
	Hind tarsi dark to reddish
161	Propodeum with majority of hairs black, at most with scattered pale hairs
	162
_	Propodeum with majority of hairs pale, at most with scattered dark hairs
	163
162	S8 apically emarginate. Slightly smaller, 9–11 mmpraecox (Scopoli)
_	S8 apically truncate. Slightly larger, 10–12 mmapicata Smith
163	Basal mandibular tooth short. Apical margins of S2-4 with long loose hairs
	that do not form clear fringes, hairs longer than the length of the hind basi-
	tarsis. Restricted to northern Spain. Flying later in the year (June-August),
	associated with shrubs, particularly Rubus (Rosaceae)fucata Smith
_	Basal mandibular tooth long. Apical margins of S2-4 with dense fringes
	composed of short hairs, these hairs not exceeding the length of the hind
	basitarsis. Flying earlier in the year (March-June)164
164	Hind tarsi reddish. Genital capsule more elongate, gonocoxal teeth com-
	paratively weak. Restricted to areas around the Pyrenees and Cantabrian
	Mountains. Flying during March-May, associated with Salix (Salicaceae)
_	Hind tarsi dark. Genital capsule more compact, gonocoxal teeth strongly
	produced. More widespread across northern Iberia into northern Portugal.
	Flying during May-June, associated with <i>Vaccinium</i> (Ericaceae)
	Letterstead

165	Mandible without any kind of angulation at its base. Mesosoma with bright
	reddish brown hairs, with some black hairs on the mesepisternum. Associ-
	ated with Salix (Salicaceae)
_	Mandible at its base with an angulation. Mesosoma never with black hairs
	laterally. Associated with a wider range of flowering trees and shrubs166
166	Mandible at its base with angulation forming a strong 90° angle. In fresh
	specimens, clypeus with golden hairs. Terga with marginal areas normal,
	marginal area of T3 occupying at most 30% of segment length. Restrict-
	ed to the Pyrenees with an isolated population in the Sierra de Cazorla
	(Jáen)
_	Mandible at its base with angulation rounded, forming an obtuse angle (c. 120°).
	In fresh specimens, clypeus with white hairs. Terga with marginal areas broad,
	marginal area of T3 occupying 60% of segment length. More widespread across
	central and northern Iberia into northern Portugal synadelpha Perkins
167 (147)	Measured along ventral margin, A3 twice as long as A4
_	Measured along ventral margin, A3 longer or shorter than A4, but never
	twice as long
168	S8 unique, elongate on narrow step, apical portion medially constricted and
100	apically strongly emarginate (Fig. 88H) paucisquama Noskiewicz
_	S8 different, not of this shape
169	Fore margin of the clypeus upturned. Propodeal triangle broad and well-de-
10)	fined by raised lateral carinae, margins extending almost to the lateral edges of
	the metanotum, internal surface covered in raised irregular carinae of a similar
	width, thus appearing regular and consistent. Pronotum with weak or strong
	humeral angle. Genital capsule, see Fig. 45E, F (subgenus <i>Ovandrena</i>) 170
	Without this combination of characters
170	Larger, 9–10 mm. Tongue with outer surface of galea clearly punctate, punc-
170	
	tures separated by 1–2 puncture diameters. Pronotum with humeral angle com-
	paratively weak, and fore margin of the clypeus only weakly upturned. Sterna
	with weak and sparse fringes on apical margins. Tergal punctation compara-
	tively larger and coarser. Genital capsule, see Fig. 45Foviventris Pérez
_	Smaller, 7–8 mm. Tongue with outer surface of galea more or less smooth
	and shiny, without obvious punctures. Pronotum with humeral angle com-
	paratively strong, and fore margin of the clypeus more strongly upturned.
	Sterna with strong and dense fringes on apical margins. Tergal punctation
	comparatively fine. Genital capsule, see Fig. 45E
171	Clypeus smooth and shiny over almost its entire surface, regularly punctate,
	punctures separated by 0.5-1 puncture diameters. Metasoma elongate, es-
	sentially parallel-sided, surface finely shagreened, more or less smooth and
	shiny, deeply punctate, punctures separated by 0.5-2 puncture diameters.
	Genital capsule with gonocoxal teeth apically diverging (Fig. 89A)
_	Without this combination of characters172

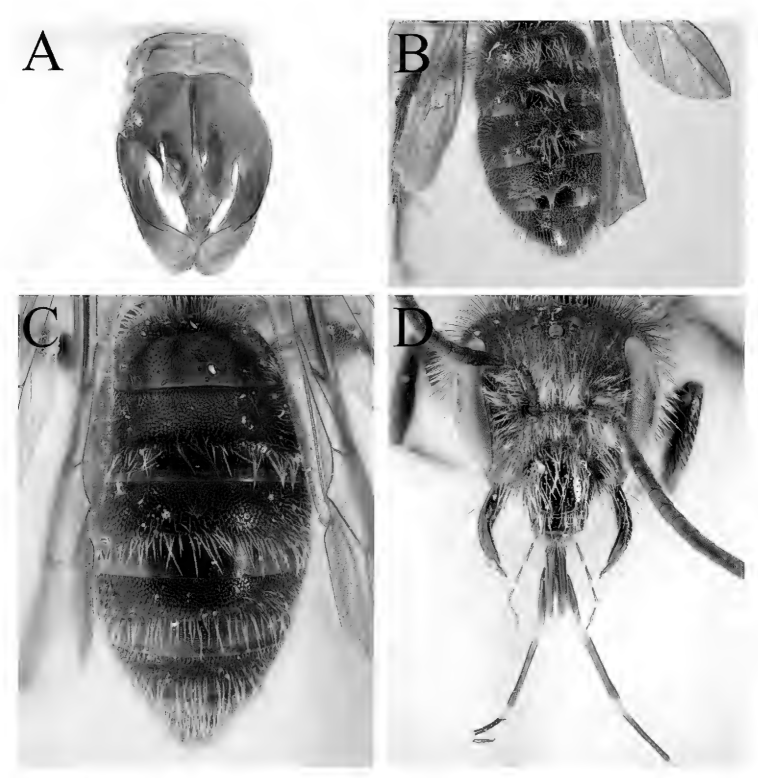


Figure 89. Andrena (incertae sedis) alluaudi Benoist, 1961, male **A** genital capsule, dorsal view; Andrena (Euandrena) fortipunctata Wood, 2021, male **B** terga, dorsal view; Andrena (Aenandrena) hystrix Schmiedeknecht, 1883, male **C** terga, dorsal view **D** Andrena (Euandrena) solenopalpa Benoist, 1945, male **D** face, frontal view.

	eyes. Terga never lightened reddish laterally. Known only from mountainous areas in north-western, central, and south-eastern Spain (Léon, Ávila,
	Jáen)
_	Tergal discs not strongly humped (Fig. 89C). Margins of T2-4 basally
	with long whitish plumose hairs that arise from the junctions with the
	discs and overly the marginal areas without obscuring the underlying sur-
	face. Facial pubescence uniformly whitish. Terga often reddish laterally,
	view in profile. Widespread across Iberia
174	Head elongate, mouthparts extremely long, twice the length of the head,
	labial palps alone equal the length of the head (Fig. 89D). Associated with
	Lithodora (Boraginaceae)solenopalpa Benoist
_	Head shorter, never with the labial palps equalling the length of the head
	175
175	Gonostyli with outer margin concave, penis valves with weak lateral hyaline
	extensions (Fig. 90A). Terga strongly microreticulate, dull. Clypeus medi-
	ally with unclear and slightly raised impunctate midline, otherwise densely
	punctate, underlying surface largely smooth and shiny. Restricted to areas
	close to the Pyrenees. Associated with Symphytum (Boraginaceae)
	symphyti Schmiedeknecht
_	Without this combination of characters176
176	A4–13 ventrally covered with shiny scales, contrasting A3 which is ventrally
	dull (c.f. Fig. 88B). Tergal margins widely lightened yellow-orange hyaline.
	Scutellum with intermixed brown-orange and black hairs. Terga very finely
	shagreened, more or less smoothly shiny, regularly punctate, punctures sepa-
	rated by 1–2 puncture diameters
_	Without this combination of characters
177	Face with pale hairs medially and black hairs along the inner margin of the
	compound eyes. Genital capsule with surface of the gonocoxae with distinc-
	tive latitudinal granular shagreen, this extending onto the basal parts of the
	gonostyli (Fig. 90B-D). Terga shagreened, with large scattered punctures
	on tergal discs, these almost becoming crater-like on T1, but without clearly
	raised rims. (former subgenus Zonandrena, partim)
_	Without this combination of characters (for species with contrasting black
	hairs along the inner margin of the compound eyes, see the next couplet)
4.50	
178	Genital capsule with outer margins of gonostyli weakly emarginate (Fig.
	90B, C)
_	Genital capsule with outer margins of gonostyli more or less straight, with-
4.50	out emargination (Figs 15B–D, 90D)
179	Penis valves comparatively broad (Fig. 90B). Rare, restricted to the Pyr-
	eneesgravida Imhoff (partim)
_	Penis valves comparatively narrow (Fig. 90C). Restricted to steppic areas in
	central Spainsoror Dours

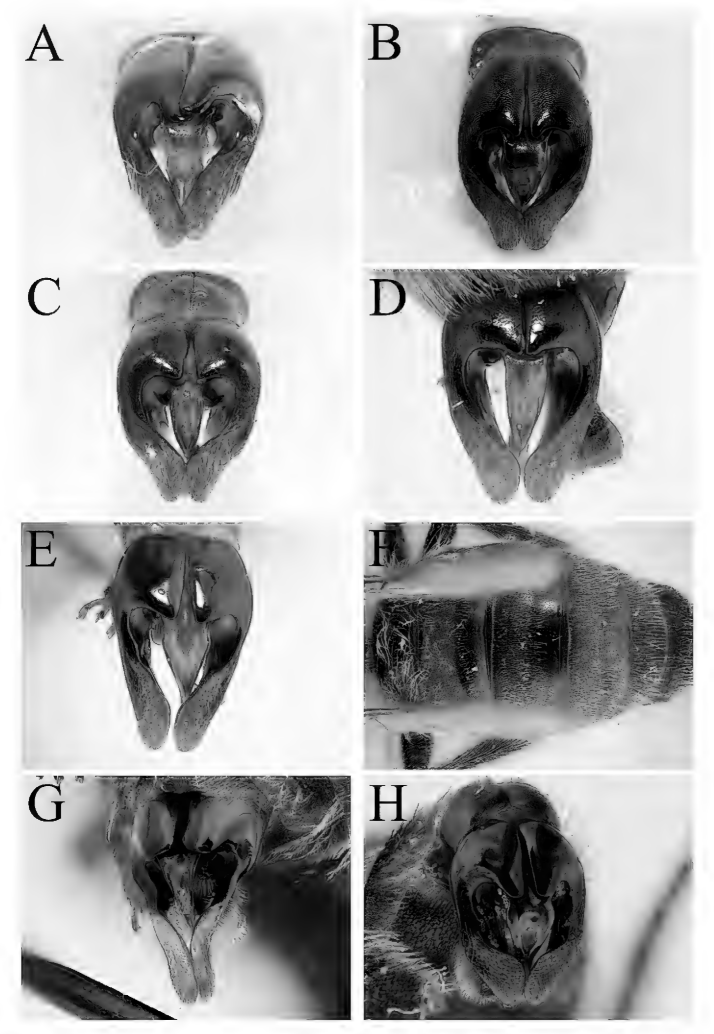


Figure 90. Andrena (Euandrena) symphyti Schmiedeknecht, 1883, male A genital capsule, dorsal view; Andrena (Melandrena) gravida Imhoff, 1832, male B genital capsule, dorsal view; Andrena (Melandrena) soror Dours, 1872, male C genital capsule, dorsal view; Andrena (Melandrena) vulcana Dours, 1873, male D genital capsule, dorsal view; Andrena (Melandrena) bicolorata (Rossi, 1790), male E genital capsule, dorsal view; F terga, dorsal view; Andrena (Lepidandrena) baetica Wood, 2020, male G genital capsule, dorsal view; Andrena (Lepidandrena) pandellei Pérez, 1895, male H genital capsule, dorsal view.

180	Genital capsule with penis valves comparatively narrow (Fig. 90D). Terga microreticulate, dull to weakly shiny. Punctures on disc of T2 less visible against background sculpture ²⁶
_	Genital capsule with penis valves comparatively broad (Fig. 15B–D). Terga shagreened, weakly shiny. Punctures on disc of T2 more clearly visible against weaker background sculpture ²⁶
181	Face predominantly white-haired, with black hairs along inner margin of compound eyes. Genital capsule elongate, relatively featureless (Fig. 90E). Tergal discs densely and finely punctate, punctures separated by 0.5 puncture diameters (Fig. 90F). Tergal margins slightly but distinctly depressed, with small, fine, and obscure punctures, the margins thus strongly contrasting the discs; underlying surface finely shagreened
	and weakly shiny
182	Tergal discs with uniformly pale pubescence (Fig. 90F) <i>bicolorata</i> (Rossi)
_	Tergal discs with intermixed black and white pubescence
	florentina Magretti
183	Terga densely and regularly punctate, punctures separated by up to 1 puncture diameter, underlying surface weakly shagreened to smooth and shiny. T2–4 with distinct white apical hair bands, often abraded and interrupted medially (remaining <i>Lepidandrena</i>)
_	Terga shagreened, with large 'crater punctures' with raised rims. Tergal margins without apical hair bands (remaining <i>Chlorandrena</i>)
184	Genital capsule elongate, gonostyli extremely long, many times longer than wide (Fig. 90G). Hind basitarsi lightened orange
_	Genital capsule with gonostyli compact, not extremely narrow and elongate (Fig. 90H). Hind basitarsi dark
185	S8 with short hairs that do not noticeably project laterally. Tarsal segment 5 of the hind leg elongate and bent. Slightly larger, 10–11 mm
_	S8 with long, laterally projecting hairs. Tarsal segment 5 of the hind leg not noticeably bent. Slightly smaller, 9–10 mm
186	Process of S8 large, with triangular-shaped lateral projections covered with projecting hair tufts (Fig. 91A; <i>taraxaci</i> -group)
_	Process of S8 large or small, but more or less parallel-sided, without triangular lateral projections or hair tufts (Fig. 91B–D)
187	Ventral surface of S8 with long, ventrally projecting hairs, clearly visible in profile. Apex of S8 emarginate
_	Ventral surface of S8 without ventrally projecting hairs. Apex of S8 rounded, never medially emarginate

²⁶ Separation of the these species is very challenging without access to confidently determined reference material.

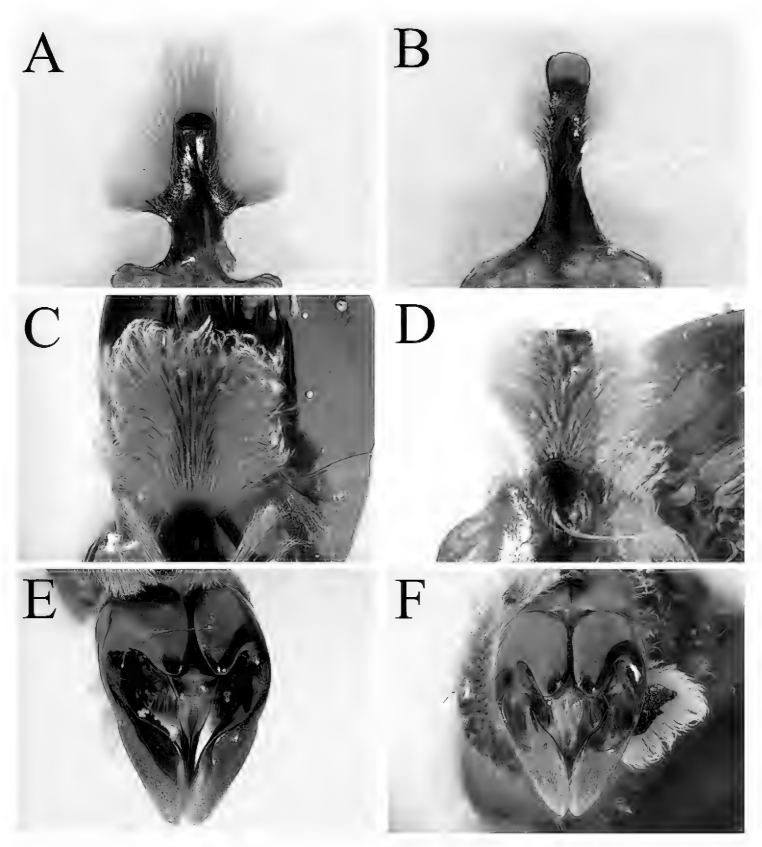


Figure 91. Andrena (Chlorandrena) rhenana Stöckhert, 1930, male **A** S8, dorsal view; Andrena (Chlorandrena) cinerea Brullé, 1832, male **B** S8, dorsal view; Andrena (Chlorandrena) livens Pérez, 1895, male **C** S8, ventral view **E** genital capsule, dorsal view; Andrena (Chlorandrena) abrupta Warncke, 1967, male **D** S8, ventral view; Andrena (Chlorandrena) agnata Warncke, 1967, male **F** genital capsule, dorsal view.

189	Process of S8 relatively short, ventral surface apically glabrous, viewed ventrally with hairs covering an area more or less as broad as long (Fig. 91B)
_	Process of S8 relatively long, ventral surface entirely covered with hairs, viewed ventrally with hairs covering an area clearly longer than broad (Fig. 91C, D)
190	Larger, 11–12 mm. S8 strongly broadened apically, here broader than the stem (Fig. 91C). Body dark, without metallic reflections
_	Smaller, 9–10 mm. S8 parallel-sided along its entire length, not apically broadened (Fig. 91D). Clypeus, scutum, scutellum, and tergal discs with obscure greasy green metallic reflections
191	Gonostyli with inner margins more or less evenly rounded (Fig. 91E). Terga comparatively more strongly shagreened, weakly shiny. Distributed throughout Iberia
_	Gonostyli with inner margins strongly flattened and produced into a raised ridge (Fig. 91F). Terga with shagreenation reduced, terga more strongly shiny in direct comparison. Rare, known only from as single specimen captured near to Madrid
192	Clypeus strongly domed, underlying surface weakly shagreened laterally, smooth and shiny over the majority of its area; clypeus largely dark, with at most occasional hints of metallic green reflections. Terga with bases very weakly depressed. Found in dry to steppic areas in central Spainelata Warncke
_	Clypeus weakly domed, underlying surface shagreened, weakly shiny; metallic green reflections present. Terga with bases strongly depressed. Found in areas close to or on the coast in southern Portugal and Spain
193 (167)	Measured along ventral margin, A3 shorter than or as long as A4194
_	Measured along ventral margin, A3 slightly longer than A4215
194	A3 extremely short relative to A4, at most ¹ / ₅ the length (Fig. 81A). Terga dark. Genital capsule distinctive, with extremely narrow parallel-sided penis valves (Fig. 81B). Rare, known from eastern Spain (Jáen, Soria, Teruel)
_	Without this combination of characters195
195	Fore margin of clypeus upturned. Gena slightly exceeding width of compound eye. T2–4 with dense white apical hair bands (Fig. 40G). Genital
	capsule compact, rounded, with gonocoxal teeth apically diverging, gono-
	styli apically broadened and flattened (Fig. 40H)
106	Without this combination of characters
196	A4–13 ventrally covered with shiny scales, contrasting A3 which is ventrally dull (Fig. 88B)
107	A4–13 not noticeably different from A3, without shiny scales200
197	Smaller, 7–8 mm. Tergal discs finely shagreened, more or less smooth and shiny, with deep and distinct punctures, punctures on disc of T1 separated

	by 1–2 puncture diameters, on discs of T2–4 separated by 0.5–1 puncture diameter. Tergal margins broadly lightened yellow-orange hyaline198
_	Larger, 10–11 mm. Tergal discs shagreened to coarsely microreticulate, at most weakly shiny, with fine regular punctures or coarse 'crater punctures'. Tergal margins at most with their apical rims narrowly lightened hyaline brown
198	Clypeus, scutum, and scutellum laterally shagreened, medially smooth and shiny (Fig. 88B). Restricted to temperate areas in northern Portugal and
_	Spain
	across Iberia
199	Terga shagreened, weakly shiny, with fine and dense punctation. Tergal margins comparatively weakly depressed, with narrow tight apical white hair bands. Hind basitarsi dark. Throughout Iberia
_	Terga strongly microreticulate, with coarse 'crater punctures'. Tergal margins comparatively strongly depressed, with long loose white apical hair bands that exceed the length of the marginal areas. Hind basitarsi lightened orange. Restricted to high altitude sites (>1200 m) in the Pyrenees
200	Larger species, 12–14 mm, usually with abundant black, white, or brown pubescence, usually with entirely dark black facial hair. Ocelloccipital distance >2 times the diameter of a lateral ocellus. Terga always without apical hair bands, sometimes with lateral hair patches (<i>Melandrena</i> partim)201
_	Smaller species, 9–10 mm, usually with subdued pubescence, facial hairs often pale, though dark in <i>A. lepida</i> and <i>A. propinqua</i> . Ocelloccipital distance often shorter, <2 times the diameter of a lateral ocellus. Terga with or without hair bands
201	Face and mesepisternum with long white pubescence, mesosoma dorsally with bright light brown pubescence. Univoltine, flying April-June. Restricted to temperate parts of northern Spain ²⁷
_	Without this pattern of pubescence; face either with extensive dark pubescence, or mesepisternum with dark pubescence, or mesosoma dorsally with dark pubescence
202	Body with only black and white pubescence203
_	Body with at least some brown pubescence

²⁷ Note, separation of *Melandrena* species in this group is usually made on the basis of colouration. However, these taxa are extremely variable, and certain characters that work in elsewhere in Europe do not work in Iberia. Caution should be taken; powerful genetic techniques are required to robustly delineate these species in an Iberian context.

203	Body typically more extensively dark haired, face and mesepisternum normally with entirely black hairs. T2–4 laterally with strongly contrasting
	dense patches of white pubescence ²⁸ albopunctata (Rossi)
_	Body often with extensive pale hairs, face sometimes white-haired with
	black hairs laterally, sometimes entirely black-haired. Terga often entirely
	black haired, laterally without dense patches of white pubescence. In pale
	forms, if T2-4 have loose white hair fringes laterally, then mesepisternum
	always white-haired ²⁸ morio Brullé
204	Terga shagreened, obscurely punctate, with punctures disappearing into the
	underlying sculpture
_	Terga at most weakly shagreened, sometimes smooth and shiny, at least
	weakly shiny. Discs of T2-5 clearly and usually densely punctate, punctures
	typically separated by 1 puncture diameter205
205	Marginal areas of terga with narrow section of apical rim lightened hyaline-
	brown, tergal discs with weak bronzy reflections. Face with mixture of dark
	brown and black hairs
_	Marginal areas of terga with apical rim dark, not lightened, tergal discs dark,
	without weak bronzy reflections. Face with uniformly black hairs206
206	Disc of T1 with dense punctures, punctures separated by up to 2 punctures
	diameters ²⁹
_	Disc of T1 with more scattered punctures, punctures usually separated by
	over 3 puncture diameters ²⁹ <i>thoracica</i> (Fabricius)
207	A3 extremely short, at most 0.5 times the length of A4 ³⁰ 208
_	A3 longer, at least 0.8 times as long as A4, often as long as A4209
208	Facial hair almost entirely black, with a few scattered light brown hairs
	around the antennal insertions. Clypeus densely punctate, punctures sep-
	arated by 0.5 puncture diameters, underlying surface smooth and shiny.
	Widespread across Iberia
_	Facial hair intermixed light and dark. Clypeus more sparsely and irregu-
	larly punctate, punctures separated by 1-2 puncture diameters, underlying
	surface microreticulate and dull. Very rare, known only from one specimen
	from Cádiz rhypara Pérez
209	Genital capsule distinctive, elongate, basally narrowed (Fig. 92A). Tergal
	discs with punctation becoming sparser laterally, most clearly visible on T2
	210
_	Genital capsule otherwise. Tergal discs with uniform punctation211

²⁸ These two species can be tricky to separate due to large colour variation within *A. morio* following the synonymy of the more extensively pale-haired *A. hispania*. Association with females should be made.

²⁹ These two species are very difficult to separate in the male sex, and positive determination is not possible in many cases.

³⁰ Note, male Simandrena are very difficult to identify; association should be made with females.

210	Face with predominantly dark hairs, with some light hairs intermixed around the antennal insertions. Scutum polished, shiny. Hind tarsi dark.
	Common throughout Iberia
_	Face with entirely light hairs. Scutum shagreened and dull. Hind tarsi light-
	ened orange. Restricted to temperate areas in northern Spain
211	Face with entirely bright pubescence, without any dark hairs laterally. Terga
211	entirely smooth and shiny, without shagreenation combinata (Christ)
_	Face with at least some dark hairs laterally. Terga often with shagreenation.
	212
212	Terga entirely shagreened and dull to weakly shiny at most. Clypeus with
	fine granular shagreen, relatively shallowly punctate, medially with weak
	impunctate longitudinal midline. Discs of T2–4 densely punctate, punc-
	tures separated by 0.5 puncture diameters
_	Terga less strongly shagreened, weakly to strongly shiny. Structure of clypeus
	variable, from shagreened to smooth and shiny, more strongly and deeply
	punctate. Terga less densely punctate, punctures on discs of T2-4 separated
	by at least 1 puncture diameter
213	Mesepisternum and propodeum with abundant black-brown hairs; remain-
	ing pubescence whitish. Scutum comparatively less strongly shagreened,
	weakly shiny. Restricted to areas close to the Pyreneesthomsonii Ducke
_	Mesepisternum, dorsum of mesosoma, and propodeum with bright yellow-
	ish pubescence, at most with occasional scattered black hairs. Scutum com-
	paratively more strongly shagreened, dull214
214	Clypeus between the punctures smooth and shiny. Tergal discs without sha-
	green, smooth and shiny, clearly visible at the base of T2. More widespread
	across Iberia ³¹
_	Clypeus between the punctures shagreened, at most weakly shiny. Tergal discs
	shagreened, weakly shiny, most clearly visible on the base of T2. Restricted to
	the Pyrenees and Cantabrian Mountains ³¹
215 (193)	Genital capsule unique within the Iberian fauna (Fig. 92B). Restricted to the
	Pyrenees and Cantabrian Mountains
_	Genital capsule otherwise
216	Head elongate, only slightly wider than long. Clypeus shagreened and dull in
	its basal half, polished and shiny in its apical half (Fig. 54C). Mesepisternum
	laterally with long strongly plumose light brown hairs (Fig. 54A). Genital
	capsule simple, with penis valves progressively narrowing apically (Fig. 54F).
	Known only from south-western Spain (Cádiz, Sevilla), flying in the very
	early spring (January–March) ramosa Wood
_	Combination of characters otherwise217

³¹ These two species are challenging to separate, and there may be taxonomic complexity in Iberia as well as across Europe. Further study is required.

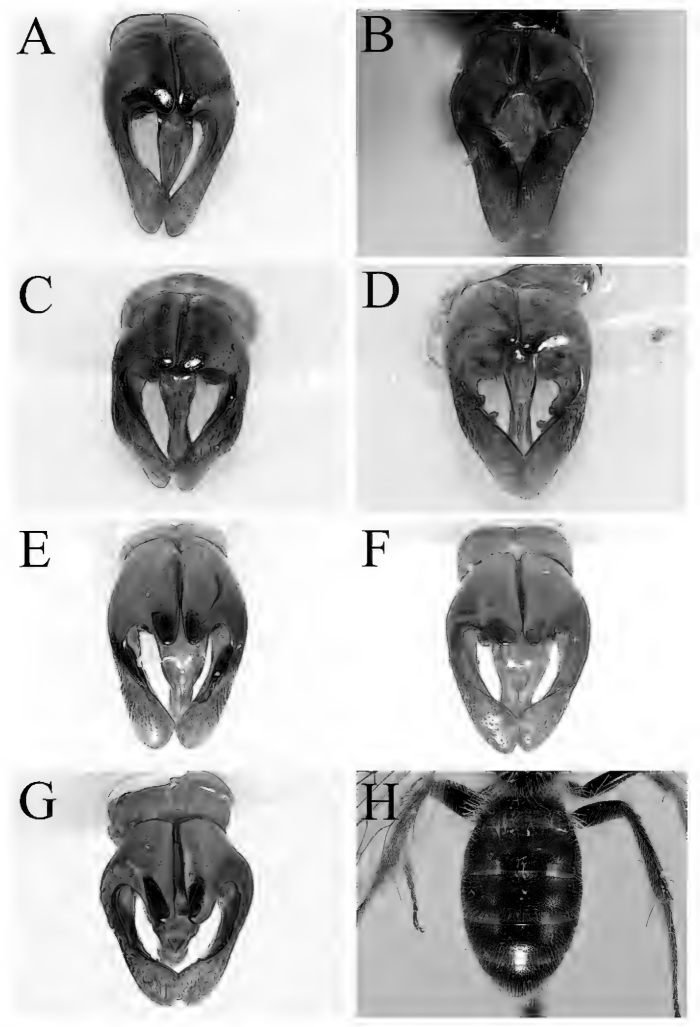


Figure 92. Andrena (Simandrena) propinqua Schenck, 1853, male **A** genital capsule, dorsal view; Andrena (Ulandrena) polita Smith, 1847, male **B** genital capsule, dorsal view; Andrena (Euandrena) granulosa Pérez, 1902, male **C** genital capsule, dorsal view **H** terga, dorsal view; Andrena (Euandrena) bicolor Fabricius, 1775, male **D** genital capsule, dorsal view; Andrena (incertae sedis) corax Warncke, 1975, male **E** genital capsule, dorsal view; Andrena (Leucandrena) parviceps Kriechbaumer, 1873, male **F** genital capsule, dorsal view; Andrena (Leucandrena) argentata Smith, 1844, male **G** genital capsule, dorsal view.

217	Clypeus domed, covered with dense network of coarse and strongly raised transverse wrinkles. Dorsolateral surface of propodeum with dense network of raised rugosity that is almost indistinguishable from the propodeal triangle, this network extending onto the lateral faces of the propodeum. Terga
	smooth and shiny with regular deep punctures, without a hint of shagreenation
_	Without this combination of characters218
218	Tergal margins distinctly depressed, depressions medially occupying 40% of the visible segment (Fig. 89C). Tergal discs densely punctate, punctures separated by 0.5–1 puncture diameters; tergal margins impunctate, lightened hyaline brown, strongly contrasting tergal discs. Margins of T2–4 basally with long whitish plumose hairs that arise from the junctions with the discs and overly the marginal areas without obscuring the underlying surface.
	Facial pubescence uniformly whitish. Terga often reddish laterally, view in
	profile
_	Without this combination of characters219
219	Penis valves narrow, more or less parallel-sided along their length (Fig. 92C, D; <i>Didonia</i> and remaining <i>Euandrena</i>)
_	Penis valves basally broadened, clearly wider basally than medially or apically (Fig. 92E–G)
220	Head relatively elongate. Process of the labrum large, slightly wider than long. Clypeus domed, with large irregular punctures, punctures separated by 0.5–2 puncture diameters, with unclear impunctate longitudinal midline. In fresh specimens, terga with clear unbroken apical hair fringes. Bivoltine, March-April and again in June-July
_	Head less elongate. Process of the labrum smaller, more clearly wider than long. Clypeus more weakly domed, shiny to shagreened. Terga without apical hair fringes
221	Apical margins of T1–5 widely lightened hyaline-yellow (Fig. 92H). Terga impressed basally and apically, thus tergal discs appear humped. Species associated with Cistaceae
_	Tergal margins at most with apical margins narrowly lightened yellowish. Tergal discs comparatively flat. Species not strongly associated with Cistaceae 223
222	Base of terga and tergal margins comparatively strongly impressed, margins clearly separated from disc by a visible 'step' (Fig. 92H). Tergal discs coarsely and densely punctate, punctures separated by 2 puncture diameters ³²
_	Base of terga and tergal margins comparatively weakly impressed, margins not clearly separated from discs by a visible 'step', almost level with discs medially. Tergal discs less densely punctate, punctures separated by 3–4 puncture diameters ³² <i>vulpecula</i> Kriechbaumer

³² These two species are difficult to separate morphologically. Comparison with confidently determined reference material should be made.

223	Gonostyli apically truncate, outer margin forming an acute point. A3 only slightly longer than A4. Pubescence entirely light brown with exception of
	occasional dark hairs laterally on the face. Restricted to temperate areas close to the Pyrenees
_	Gonostyli with outer margin rounded, never forming an acute point. A3
	visibly longer than A4224
224	Light hairs on dorsal parts of mesosoma, terga and sterna yellowish white
	in fresh specimens. Erect hairs on disc of T4 usually predominantly dark.
	Facial pubescence often entirely dark. Bivoltine, can be recorded between
	March and August. Widespread across Iberia 33 bicolor Fabricius sensu
	lato (two mitochondrial lineages are present in A. bicolor; to date, only the
	southern lineage has been found in Iberia)
_	Light hairs on dorsal parts of mesosoma, terga and sterna snow white, with-
	out yellowish hue even in very fresh specimens. Disc of T4 at most with a few
	isolated, erect dark hairs. Face always with some grey-white hairs medially, at
	least between and around antennal sockets. Univoltine, April-June. Restricted
	to mountainous areas in northern and north-western Spain ³³ allosa Warncke
	(note, the Spanish taxon may be distinct from populations in Central Europe)
225	Face medially with white hairs, laterally with clear line of black hairs along
	the inner margin of the clypeus. Genital capsule with outer margins of gon-
	ostyli weakly emarginate. Rare, restricted to the Pyrenees
	gravida Imhoff (partim)
_	Face entirely pale-haired, at most with scattered black hairs laterally. Genital
	capsule with outer margins of gonostyli straight, without emargination 226
226	Terga very weakly shagreened, more or less smooth and shiny. Tergal discs
	with extremely scattered punctures, punctures separated by 3-4 puncture
	diameters or more. Genital capsule, see Fig. 92F
_	Terga either more strongly microreticulate (at least on discs of T2-3) or
	clearly and densely punctate, punctures separated by 1-2 puncture diam-
	eters
227	Tergal discs finely microreticulate, with obscure and scattered punctures,
	punctures separated by 2-4 puncture diameters. Tergal margins with narrow
	broadly interrupted hair fringes apicallybarbilabris (Kirby)
_	Tergal discs at most finely shagreened, more or less smooth and shiny, clear-
	ly and regularly punctate, punctures separated by 1-2 puncture diameters.
	Tergal margins with narrow interrupted or broad uninterrupted hair bands
	apically
228	Tergal discs with punctures slightly sparser, separated by 2 puncture diam-
	eters. T2-4 apically with broad uninterrupted apical white hair bands, these

³³ Note, the Spanish male of *A. allosa* is unknown, therefore this couplet is based on the key of Praz et al. (2019) for the Alps.

Discussion

At 228 species, the Iberian fauna is slightly larger than those of Greece (c. 220) and Israel (c. 220), though the fauna of Israel is likely to eventually be larger than that of Iberia following ongoing revisions (Pisanty et al. 2022a). Within the Old World fauna, it is much smaller than that of Turkey which has an estimated 376 species (Wood 2023b), but this is less surprising as Turkey is larger (783,000 km² versus 583,000 km²), sits at the interface of Europe and Asia, hosts an even greater diversity of habitats, and is likely at the centre of diversity for this genus (Pisanty et al. 2022b). Iberia nevertheless hosts one of the top five largest *Andrena* faunas globally.

There are a number of outstanding problems that remain to be resolved, in addition to those highlighted above. Members of the subgenus Avandrena Warncke remain difficult to interpret due to their morphological variation and rarity in collections. Andrena (Avandrena) avara Warncke, 1967 sensu stricto was described from Morocco and also occurs in southern Iberia. Two additional subspecies were described from Spain, A. avara gavia Warncke, 1974 (locus typicus Madrid in central Spain) and A. avara liturata Warncke, 1974 (locus typicus Sierra de Guadarrama in central Spain), as well as two additional subspecies from North Africa. These may all represent valid species; molecular revision of this subgenus across the West Palaearctic is necessary to define species boundaries, and also to establish whether members of the Avandrena that lack spines on the posterior face of the hind femur truly belong here (Pisanty et al. 2022b). Many other groups have subspecies which Warncke described from Iberia, most pertinently A. (Micrandrena) pandosa trigona Warncke, 1975, A. (Notandrena) langadensis albipila Warncke, 1967, and A. (Truncandrena) medeninensis donata Warncke, 1967. These may also represent valid species, but without genetic data it is unwise to raise them to species status, as the results presented here illustrate that the relationships between Iberia, North Africa, and the rest of continental Europe are not always straightforward.

A total of 33 of the 228 *Andrena* species recorded here are endemic to the peninsula (14.5%). Twenty-two of the species are true Spanish endemics (9.6% endemic), whereas no species are endemic to Portugal. Of these 33 species, 31 of them were described after 1967 in just a handful of publications by a very limited number of

authors (Warncke 1967, 1975a; Wood et al. 2020a, 2021, 2022a; Wood and Ortiz-Sánchez 2022; Wood 2022; the current work). Only the endemic A. pruinosa and A. cyanomicans (see Kratochwil 2021) were described earlier (Erichson 1835; Pérez 1895). The fact that so many of Iberia's endemic species have only been described recently makes it difficult to conclude on a likely final pattern of richness. As our ability to distinguish between cryptic taxa has increased, so has our ability to quickly travel to areas with unusual climates, botanical communities, or biogeographical histories. A defining characteristic of bees of the genus Andrena is their ability to very rapidly adapt their behaviours to local conditions over evolutionary time, be that in the use of a novel or locally abundant host plant, changing their emergence date or voltinism, or surviving in drier, wetter, hotter, or colder environments. This ability likely underpins their extremely high speciation rate (Bossert et al. 2022; McLaughlin et al. 2022). For example, species described from Iberia in just the last few years include those restricted to saline soils (Andrena juliana), a species which uses generic Brassicaceae species but which flies at the unusual time of December (Andrena gades), an early emerging taxon restricted to south-western Spain which has avoided capture until very recently due to its flight period of January-March (Andrena ramosa), a specialist of late-flowering Apiaceae which flies during September-October (Andrena foeniculae), and specialists of little-used botanical families such as Crassulaceae and Geraniaceae (Andrena omnilaevis and Andrena erodiorum). Altogether, this means that until all unusual habitats and mountainous areas have been thoroughly searched across the entirety of the bee season, new species are almost guaranteed to continue to be found.

It is possible however to comment more broadly on the biogeography of Iberian bees and notable distributional patterns. Warncke (1975a) identified four major biogeographic zones on the peninsula (excluding the Balearic Islands); a) north-western and northern Iberia from Coimbra to the Pyrenees, this area hosting Central European species, b) the northern meseta (Meseta Norte), or "Old Castille" (formerly known as Castilla la Vieja), the area of elevated open habitats north of the Sistema Central to the Cantabrian range, c) the southern meseta (Meseta Sur), or "New Castille" (formerly known as Castilla la Nueva) to the Ebro Valley, the lower area of open habitats southeast of the Sistema Central to the Ebro Valley, separated by the Sistema Ibérico, and d) the coastal zone from Coimbra in the west to Cádiz in the south up to Barcelona in the north-east, this area possessing a subtropical climate with hot summers and hosting Mediterranean taxa. These categorisations are broadly correct but do not fully capture the diversity of Iberian habitats and some of the patterns of endemism and micro-endemism. As a corollary, Warncke (1975a) writes that in the driest and warmest parts of Iberia some species have evolved into their own subspecies. It is worthwhile to highlight these areas in light of taxonomic developments in the intervening years.

The most substantial change since Warncke's (1975a, 1976) Iberian revision is the study of mountain refugia and their associated fauna. The Serranía de Ronda/Sierra de las Nieves and Sierra Nevada all host endemic *Andrena* species that have diverged from Iberian or from European/North African species (*A. contracta*, *A. ghisbaini*, *A. isolata*, and *A. ortizi*). The Sistema Central to the mountains in north-western Iberia (Serra do

Gerês, Cantabrian Mountains, etc) host endemic species that prefer cooler temperatures and are associated with herbaceous plants on acidic soil (*A. benoisti*, *A. gredana*, *A. omnilaevis*). In eastern Spain, the mountain ranges of the Sistema Ibérico, Sierra de Cazorla, and smaller ranges are understudied and support distinct species (*A. levante*) and subspecies whose status should be further investigated (*A. hattorfiana nigricauda* Wood, 2021). Because the eastern mountain ranges of Spain are somewhat isolated latitudinally, with intervening low elevation areas, it is likely that many of the species present here at elevation have experienced interrupted gene flow in their recent evolutionary history (e.g. subgenus *Taeniandrena*, Wood et al. 2021; Praz et al. 2022). The links between these montane populations should be investigated to establish their status. Members of the *A. intermedia* Thomson, 1870 aggregate may well be distinct, and populations present on the Sierra Nevada -> Sierra de Cazorla -> Sistema Ibérico -> Pyrenees may each represent a different species. Whether this pattern hold true for other groups requires further study.

The elevated plateau of central Spain clearly hosts relictual *Andrena* faunal elements. This is most clearly seen for two subgenera, *Nobandrena* Warncke, 1968 and *Parandrenella* Popov, 1958. *Nobandrena* consists of 10 species from Central Europe to Central Asia (western limit Switzerland), with one species endemic to central Spain (*A. funerea* Warncke, 1975), predominantly in the provinces of Ávila, Madrid, Salamanca, and Segovia. Likewise, *Parandrenella* consists of nine species from eastern Central Europe to Central Asia and Pakistan (European western limit is Slovenia, with an additional species in north-western Africa from Morocco to Tunisia; see Scheuchl et al. 2011, also Wood et al. 2020b), with one species endemic to central Spain (*A. taxana* Warncke, 1975), known only from the provinces of Cuenca, Madrid, and Soria. *Andrena funerea* and *A. taxana* therefore represent what must have been a broader distribution for these subgenera prior to the Quaternary period, both subgenera having estimated crown ages of around 10 million years (Pisanty et al. 2022b).

More broadly, Iberia hosts several predominantly eastern species in steppic or dry areas, such as A. lateralis (Spain and Portugal, southern Balkans to the Central Asia), A. (Melandrena) soror Dours, 1872 (Spain, Morocco, Turkey), and A. urdula (Spain, Morocco, and Greece). Iberia has strong faunal links to Morocco, and particularly the Middle Atlas. Due to the nature of European mountain chains, outside of Iberia, raised steppic areas are rare or cover only very small areas. In Central Spain, Morocco, and Turkey, extensive raised areas (the *mesetas* of Old and New Castille) of steppic habitat can be found. This elevated steppe differs from the Great Eurasian Steppe that runs from the Pannonian basin (predominantly eastern Austria, Hungary, southern Slovakia, western Romania, northern Serbia) to Mongolia and northern China, and hosts a fauna that supports the same evolutionary lineages (e.g. Nobandrena and Parandrenella), but often contains different species, many of which are endemic. This link between Iberia and Morocco can be seen particularly strongly in the Middle Atlas. In addition to the finding of *A. relata* in the Middle Atlas (Wood et al. 2020b), another species previously considered to be endemic to Spain, A. nebularia, can be found on the high steppe in the eastern Middle Atlas. This link to the Middle Atlas

can also be seen on the Sierra Nevada for *A. isolata* and *A. ortizi* which are genetically most closely related to what are undescribed species in the Middle Atlas. Additional genetic work and further surveys in the Middle Atlas are likely to strengthen these links further, as well as further surveys on the Sierra Nevada and the mountains of southern Spain.

The extraordinary nature of central Spain and its rich Andrena fauna is well-illustrated by both historical (particularly those of Dusmet that were revised by Warncke 1975a, 1976) and recent collections. During a single eight-day period from the 13th to 19th of May 2021 in the provinces of Ávila, Guadalajara, Madrid, Segovia, and Toledo I collected a total of 70 Andrena species, including the Iberian endemic A. baetica, A. benoisti, A. corax, A. elata, A. funerea, A. lecana, A. murana, A. parata, A. pruinosa, and A. varuga and also the restricted A. lateralis, A. monilia, A. nebularia, A. soror, A. relata, and A. urdula. Whilst large parts of the Sistema Central mountain range are protected as nature reserves or regional parks, the same is not true for the steppe habitats at lower altitudes, both north and south of this mountain range. Most collecting sites were habitat fragments sandwiched between urban development or intensive agriculture, though some formed a mosaic with land under a lower-intensity management regime. The areas south of Madrid that were so extensively collected by Dusmet and which represent the *loci typici* for many of the species described by Warncke have either been lost to urban development or are threatened by its encroachment (e.g. Andrena montarca was described from Montarco which is now a suburban park). Given the endemic nature of this fauna, these areas are deserving of a greater level of protection than they currently receive.

Away from central Spain, there are more obvious links between southern Spain and the North African Andrena fauna. Warncke (1976) recorded three predominantly North African taxa in the extreme south of Spain, specifically A. (Notandrena) microthorax Pérez, 1895 (Cádiz), A. (Truncandrena) minapalumboi Gribodo, 1894 (Cádiz, Alicante), and A. rhypara (Cádiz). I have seen no other material of these species, and their current status in Spain is unclear; they may be locally extinct. In contrast, recent workers have discovered three North African species in southern Spain, namely A. (Truncandrena) varia Pérez, 1895 (Córdoba; Ortiz-Sánchez 2020), A. laurivora (Huelva and Sevilla; Wood et al. 2021), and A. melacana (Albacete, Cádiz, Granada, Málaga; Wood and Ortiz-Sánchez 2022). Given the lack of historical and contemporary collecting in southern Spain, it is very difficult to answer whether or not these species were always present, whether the newly detected species are recent arrivals, or whether there is a constant turnover of North African species in southern Spain that regularly colonise and then become locally extinct. Increased recording of Andrena specifically and Iberian bee species more broadly is required in order to better understand faunal interchanges across the Strait of Gibraltar.

Finally, the difference between the size of the Portuguese (128) and Spanish (228) *Andrena* faunas is large and notable. Though Portugal has a higher density of *Andrena* species due to its much smaller size, it hosts no endemic *Andrena* species compared to the 22 species endemic to mainland Spain. As highlighted above, the Portuguese

Andrena fauna is so much smaller due to the almost complete absence of high northern mountains and their associated Euro-Siberian fauna (the Serra da Estrella reaches to 1,993 m but is isolated from the high Cantabrian Mountains of northern Spain), the lack of elevated steppe (limited to north-eastern Portugal around Almeida or the Douro valley), the lack of high mountains in the south (the Sierra de las Nieves reaches 1,919 m and the Sierra Nevada 3,479 m), and finally the absence of the very hot and dry Mediterranean habitat from Cádiz to Alicante and Valencia that hosts both North African species as well as restricted endemic species. When taken collectively, it is this enormous variety of habitats, isolated mountains, and Atlantic and Mediterranean influences that has shaped and generated the rich Iberian Andrena fauna, and continues to provide taxonomic surprises and ecological delights.

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Supplementary material I

Iberian Andrena species checklist

Authors: Thomas J. Wood Data type: National checklist

Explanation note: Iberian *Andrena* species checklist, with national totals for Portugal and Spain, details on *loci typici* for species described from Iberia, and dietary niche classifications.

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